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GenCore version 5.1.4_p5_4578

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OM protein - protein search, using sw model

Run on: March 13, 2003, 19:10:37 ; Search time 18 Seconds (without alignments)

85.453 Million cell updates/sec
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Title:
Perfect score: 84
Sequence:
1 NNATEYFKIDNVKAR 16
Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5
Searched: 283224 segs, 96134422 residues

Total number of hits satisfying chosen parameters:
Minimum DB seq.length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
Database: PIR\_73:\*
1: pir1:\*
2: pir2:\*
3: pir3:\*
4: pir4:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

kininogen, LAW pre K-kininogen, HAW pre K-kininogen, LAW I Kininogen, HAW I Kininogen, HAW II Major acute phase T-kininogen I prec major acute phase T-kininogen, LAW I Kininogen, HAW I Kininoge hypothetical prote hypothetical prote protein F3M18.12 [ hypothetical prote Nuc.2 protein - Ne TPR-repeat contain hypothetical prote ICP18.5 homolog -hypothetical prote hypothetical prote hypothetical prote hypothetical prote conserved hypothet prote hypothetical phypothetical phy Description SUMMARIES KGBOH1 B97948 AH1213 B84791 F69403 G75105 T23191 F86410 T32285 S72479 KGHUH1 A28055 A25486 KGBOL2 KGRDHZ KGRTM KGRTT1 A23897 B28055 DB Query Match Length 895 996 1007 1066 346 8899988888888444 4477889888888888888 Score Result Ş.

hypothetical prote	hypothetical prote	probable tail comp	uncharacterized pr	hypothetical prote	hypothetical prote	hypothetical prote	hypothetical prote	Zn-dependent hydro	hypothetical coile	hypothetical prote					ō,
H85035	E90733	685583	в97033	D90132	A98248	F97339	T28070	E97075	T38834	D87637	B84996	T25830	T31994	A89769	F82150
~	N	~	~	7	~	~	~	7	~	~	ď	N	~	~	7
532	1021	1021	2817	72	180	184	237	249	306	381	406	630	758	1050	224
47.6	47.6	47.6	47.6	46.4	46.4	46.4	46.4	46.4	46.4	46.4	46.4	46.4	46.4	46.4	45.8
40	40	40	40	39	39	39	39	39	39	39	39	39	39	39	38.5
30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	4.5
	532 2 H85035	532 2 H85035 1021 2 E90733 1	532 2 H85035 1021 2 E90733 1021 2 G85583	532 2 H85035 1021 2 E90733 1021 2 E95783 2817 2 B97033	47.6     532     2     H85035     hypothetical       47.6     1021     2     B99033     hypothetical       47.6     2817     2     B97033     uncharacterizal       46.4     72     2     D90132     hypothetical	47.6     532     2 H85035     hypothetical       47.6     1021     2 E90733     hypothetical       47.6     1021     2 G85583     probable tail       47.6     2817     2 B97033     uncharacterizal       46.4     72     2 D990132     hypothetical       46.4     180     2 A99248     hypothetical	47.6     532     2     H85035     hypothetical       47.6     1021     2     B90733     hypothetical       47.6     1021     2     85583     probable tail       47.6     2817     2     B97033     uncharacteriz       46.4     72     2     B90132     hypothetical       46.4     180     2     849848     hypothetical       46.4     184     2     F97339     hypothetical	47.6     532     2     H85035     hypothetical       47.6     1021     2     B90733     hypothetical       47.6     1021     2     B97033     probable tail       46.4     72     2     D90132     hypothetical       46.4     180     2     A98248     hypothetical       46.4     184     2     F97339     hypothetical       46.4     237     2     7280770     hypothetical	532 2 H85035 hypothetical 1021 2 E90733 hypothetical 1021 2 G85583 hypothetical 2817 2 B97033 nocharacteriz 72 2 D90132 hypothetical 180 2 A98248 hypothetical 184 2 F97339 hypothetical 237 2 T28070 hypothetical 249 2 E97075 Zn-dependent	47.6     532     2     H85035     hypothetical       47.6     1021     2     B90733     hypothetical       47.6     2817     2     B97033     uncharacterizal       46.4     72     2     B9132     hypothetical       46.4     180     2     A98248     hypothetical       46.4     184     2     P97339     hypothetical       46.4     237     2     T28070     zn-28070       46.4     249     2     T38834     hypothetical       46.4     306     2     T38834     hypothetical	47.6     532     2     H85035     hypochetical       47.6     1021     2     890733     hypochetical       47.6     1021     2     89583     uncharacterizal       46.4     72     2     990132     hypochetical       46.4     180     2     849848     hypochetical       46.4     237     2     728070     pypochetical       46.4     237     2     728070     zogendent       46.4     306     2     738834     hypothetical       46.4     310     2     738834     hypothetical       46.4     310     2     738834     hypothetical       46.4     310     2     738834     hypothetical	47.6     532     2 H85035     hypothetical       47.6     1021     2 E90733     hypothetical       47.6     1021     2 G85583     probable tail       47.6     2817     2 B97033     uncharacterizal       46.4     180     2 A98248     hypothetical       46.4     184     2 P97339     hypothetical       46.4     237     2 T28070     hypothetical       46.4     249     2 E97075     Zn-dependent       46.4     306     2 T38834     hypothetical       46.4     381     2 D87637     hypothetical       46.4     381     2 D87637     hypothetical       46.4     36     2 R38934     hypothetical       46.4     406     2 B87637     hypothetical       46.4     406     2 B89966     hflk protein	47.6         532         2         H85035         hypothetical           47.6         1021         2         B90733         hypothetical           47.6         2817         2         B97033         probable tail           46.4         72         2         B90132         hypothetical           46.4         184         2         A98248         hypothetical           46.4         184         2         P97339         hypothetical           46.4         237         2         P3075         Zn-Gependent           46.4         306         2         T38834         hypothetical           46.4         406         2         B84996         hflk protein           46.4         406         2         B84996         hypothetical	47.6         532         2         H85035         hypothetical           47.6         1021         2         B90733         hypothetical           47.6         2817         2         B97033         probable tail           46.4         72         2         B97033         hypothetical           46.4         180         2         A98248         hypothetical           46.4         184         2         F97339         hypothetical           46.4         237         2         T28070         Zn-6pendent           46.4         30         2         T38834         hypothetical           46.4         30         2         T38834         hypothetical           46.4         406         2         B84996         hypothetical           46.4         406         2         B84996         hypothetical           46.4         758         2         T25830         hypothetical           46.4         758         2         T35830         hypothetical	47.6 532 2 H86035 47.6 1021 2 G85583 47.6 2817 2 B97033 46.4 180 2 A99248 46.4 249 2 E97707 46.4 249 2 E97707 46.4 249 2 E97707 46.4 306 2 T38834 46.4 406 2 T38834 46.4 406 2 T38834 46.4 1050 2 T38994 46.4 1050 2 A89769

## ALIGNMENTS

	KGHULI KGHULI KAILETRATE names: alpha-2-thiol proteinase inhibitor; preprokininogen N;Alternate names: alpha-2-thiol proteinase inhibitor; preprokininogen N;Contains: bradykinin (kallidin); kininogen I; kininogen II; prokininogen C;Specias: Homo saplens (man) C;Date: 06-Jul-1982 #sequence_revision 27-Nov-1985 #text_change 08-Dec-2000 C;Accession: A01280; B25276; A27900; A27699; A31905; A34030 E;Obate: 06-Jul-1982 #seasway, T.; Shlokawa, H.; Sasaki, M. Biochemistry 23, S691-5697, 1984 A;Title: Isolation of a human cDNA for alpha-2-thiol proteinase inhibitor and its ide A;Reference number: A90490; MUID:85122621; PMID:6441591
	A; Molecule type: mRNA A; Residues: 1-427 < CHKX A; Residues: 1-427 < CHKX A; Cross-references: GB: CDES66; NID:g177889; PIDN:AAA35497.1; PID:g177890 B; Takagaki, Y.; Kitamura, N.; Nakanishi, S. J. Biol. Chem. 260, 8601-8609, 1985 A; Title: Cloning and sequence analysis of cDNAs for human high molecular weight and 1 A; Reference number: A92544; MUID:85234582; PMID:2989293 A; Accession: B25276
:	A; Molecule type: mRNA A; Residues: 1-427 < CTAK> A; Residues: 1-427 < CTAK> A; Cross-references: GB: M11437; NID:g186751; PIDN:AAB59551.1; PID:g386853 A; Cross-references: GB: M11437; NID:g186751; PIDN:AAB59551.1; PID:g386853 B; Lottspeich, F.; Kellermann, J.; Henschen, A.; Rauth, G.; Mueller-Esterl, W. in Kinins IV, part A, Greenbaum, L.M., and Margolius, H.S., eds., pp.91-95, Plenum, N A; Reference number: A27900 A; Reference number: A27900 A; Molecule type: protein A; Molecule type: protein A; Molecule type: protein A; Molecule type: Carretero, O.A.; Proud, D.; Walz, D.; Scicli, A.G. B; Mindrolu, T.; Carretero, O.A.; Proud, D.; Walz, D.; Scicli, A.G. B; Mindrolu, T.; Carretero, O.A.; Proud, D.; Walz, D.; Scicli, A.G. A; Title: A new kinin molecty in human plasma kininogens. A; Reference number: A27699; MUID:88209021; PMID:3365237
,	A; Accession: A27569 A; Molecule type: protein A; Residues: 380-389 < MIN> A; Molecule type: protein A; Residues: 380-389 < MIN> B; Macda, H.; Mateumura, Y.; Kato, H. B; Macda, H.; Mateumura, Y.; Kato, H. B; Molecule: Durification and identification of [hydroxyprolyl(3)]bradykinin in ascitic f A; Reference number: A31905; MUD:89034061; PMID:3182782 A; Accession: A31905 A; Molecule type: protein A; Residues: 381-389 < MAE> A; Residues: 381-389 < M
	KySasaguri, w.; Ireda, w.; Jedashi, w.; Arakawa, K Biochem, Biophys. Res. Commun. 150, 511-515, 1988 A;Title: Identification of [hydroxyproline(3)]-1ysyl-bradykinin released from human p A;Reference number: A34030; MUD:88106632; PMID:3337729 A;Accession: A34030 A;Wolecule type: protein A;Residues: 380-389 <sas></sas>

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A. Accession: A91153
A. Molecule type: protein
A. Residues: 379-644 < LOT-
A. Note: the bradyfuln sequence preceding the light chain sequence was not determined
B. Kellermann, J.; Lottspeich, F.; Henschen, A.; Mueller-Esterl, W.
Bur. J. Biochem. 154, 471-478, 1986
Bur. J. Biochem. 154, 471-478, 1986
A. Title: Completion of the primary structure of human high-molecular-mass kininogen.
A. Reference number: A24871; MUID:86108361; PMID:3484703
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A.Accession: A24871
A.Molecule type: protein
A.Rollecule type: protein
A.Rollecule: C.,20-380 (XEL1>
R.Kellermann, J.; Lottspeich, F.; Henschen, A.; Mueller-Esterl, W.
In Kinins IV, Greenbaum, L.M., and Margolius, H.S., ed., pp.85-89, Plenum Press, New
A.Title: Amino acid sequence of the light chain of human high molecular mass kininoge
A.Rollermore anniher: A27899
                                                                                                                                                                                        A; Molecule type: mRNA
A; Residues: 'ASSM', 233 -337 <AUE>
A; Note: differences are due to known cloning artifacts
A; Note: differences are due to known cloning artifacts
R; Lottspeich, F.; Kellermann, J.; Henschen, A.; Foertsch, B.; Muller-Esterl, W.
Eur. J. Blochem. 152, 307-314, 1985
A; Title: The amino acid sequence of the light chain of human high-molecular-mass kini
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A; Residues: 380-389 -586-586-
R; Lenarcic, B.; Gabrijelcic, D.; Rozman, B.; Drobnic-Kosorok, M.; Turk, V.
Biol. Chem. Hoppe-Seyler 369, 257-261, 1988
A; Title: Human cathepain B and cysteine proteinase inhibitors (CPIs) in inflammatory
A; Accession: S02482; MUID:89076517; PMID:3264507
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A; Molecule type: protein
A; Residues: 380-389 Amin>
A; Residues: 380-389 Amin>
A; Residues: 380-389 Amin>
A; Residues: 380-389 Amin>
A; Rato, H.; Matsumura, Y.; Kato, H.
J. Biol. Chem. 263, 16051-16054, 1988
A; Title: Purification and identification of [hydroxyprolyl(3)] bradykinin in ascitic A; Reference number: A31905; MUID:89034061; PMID:3182782
A; Accession: A31905
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A; Molecule type: protein
A; Residues: 381-389 < MAE.
B; Sasaguri, M.; Ikeda, M.; Ideishi, M.; Arakawa, K.
Biochem. Biophys. Res. Commun. 150, 511-516, 1988
A; Title: Identification of [hydroxyproline(3)]-1ysyl-bradykinin released from human
A; Reference number: A34030; MUID:88106632; PMID:3337729
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      analogues of bradykinin in
                                                             human kininogen domain 3.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    D.; Scicli, A.G.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           A; McLecule type: protein
A; McLecule type: protein
A; Residues: 1-19;189-192;310-314;381-389 < LEN1>
R; Kato, H; Matsumura, Y; Maeda, H.
FEBS Lett. 232, 252-254, 1988
A; Title: Isolation and identification of hydroxyproline analogues
A; Reference number: A61495; MUID:88211869; PMID:3366244
A; Mccession: A61495
A; Molecule type: protein
A; Residues: 380-389 < KAT1>
A; Residues: 380-389 < KAT1>
A; Accession: B61495
A; Note: this peptide had Pro-383 modified to 4-hydroxyproline
A; Robicule type: protein
A; Residues: 381-389 < KAT2>
A; Experimental source: urine
A; Rosidues: 381-389 < KAT2>
A; Residues: 381-389 < KAT2>
A; Rote: this peptide had Pro-383 modified to 4-hydroxyproline
A; Note: this peptide had Pro-383 modified to 4-hydroxyproline
A; Accession: C61495
A; Molecule type: protein
                                               A:Title: Cloning, expression and characterization of hum A; Reference number: $32422; MUID:93223854; PMID:8467916
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A; Molecule type: protein
A; Residues: 379-389, K', 390-407, O', 409-644 <KEL2>
A; Residues: 379-389, K', 390-407, O', 409-644 <KEL2>
R; Mindrolu, T.; Carretero, O.A.; Proud, D.; Walz, D.; Sc Biochem. Blophys. Res. Commun. 152, 519-526, 1988
A; Title: A new kinin molety in human plasma kininogens.
A; Reference number: A27699; MUID:88209021; PMID:3365237
              FEBS Lett. 321, 93-97, 1993
                                                                                                                                             A; Accession: S32422
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A introns: 65/3; 102/3; 131/1; 188/3; 224/3; 253/1; 310/3; 346/3; 375/3; 401/3

A introns: 65/3; 102/3; 131/1; 188/3; 224/3; 253/1; 310/3; 346/3; 375/3; 401/3

C Superfamily: Kininogen: Oystatin homology

C; Keywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyc

C; Keywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyc

C; Keywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyc

C; Keywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyc

E; 19-37/Product: LMW prokiniogen II #status predicted cMAT>

F; 19-37/Product: LMW kininogen II #status predicted cMAT>

F; 19-37/Product: LMW kininogen in #status predicted cMAT>

F; 19-37/Product: LMW kininogen in #status experimental cMDP>

F; 142-253/Domain: cystatin homology cCY2>

F; 264-375/Domain: cystatin homology cCY2>

F; 380-389/Product: LMW kininogen light chain #status experimental cACH>

F; 380-389/Product: LMW kininogen light chain #status experimental cACH>

F; 380-380/Product: LMW kininogen light chain #status experimental

F; 380-380/Cleavage site: Met-Lys (kallikrein) #status experimental

F; 387/Modified site: Het-Lys (kallikrein) #status experimental

F; 389-390/Cleavage site: Arg-Ser (kallikrein) #status experimental

F; 401/Binding site: carbohydrate (Thr) (covalent) #status absent
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Kininogen, HWW precursor [validated] - human
N;Alternate names: alpha-2-thiol proteinase inhibitor; preprokininogen; prokininogen
N;Contains: bradykinin (kallidin I); HWW kininogen I; HWW kininogen II; low molecular we
C;Species: Homo sapiens (man)
C;Date: 28-May-1986 #sequence_revision 28-May-1986 #text_change 08-Dec-2000
C;Accession: A01279; A2526; S32422; A9153; A24871; A27899; A27699; A31905; A34030; S02
R;Ohkubo, I.; Kurachi, K.; Takasawa, T.; Shiokawa, H.; Sasaki, M.
Biochemistry 23, 5691-5697, 1984
A;Title: Isolation of a human cDNA for alpha-2-thiol proteinase inhibitor and its identification of a number: A90490; MUID:85122621; PMID:6441591
                                                                                                                                                                                                                                                                                                                                                                                                                                              C.Comment: The LAWW kiningen precursor is produced from the same gene as the HAWM form (s C.Comment: Kininogen is a cysteine proteinase inhibitor, takes part in initiation of the C.Comment: Bradykinin, released from kininogen by kallikrein, is a potent vasodilator, i xyproline residue is present in the kininogen prior to the release of bradykinin.
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A.Residues: 1-389 < OHK>
A.Residues: 1-389 < OHK>
A.R.Cross-references: GB.K02566; NID:g177889
R.Takagaki, Y.; Kitamura, N.; Nakanishi, S.
J. Biol. Cham. 260, 8601-8609, 1985
A.Fitle: Cloning and sequence analysis of CDNAs for human high molecular weight and low A.Reference number: A92544; MUID:85234582; PMID:2989293
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    T.; Nakanishi, S.
                                                                                              gene and a model for its
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A;Residues: 1-592, 17,594-644 <TAK>
A;Cross-references: GB:M1437; NID:g186751; PIDN:AAB59550.1; PID:g386852
R;Auerswald, E.A.; Roessler, D.; Mentele, R.; Assfalg-Machleidt, I.
R;KItamura, N.; Kitagawa, H.; Fukushima, D.; Takagaki, Y.; Miyata, D. Blol. Chem. 260, 8610-8617, 1985
A;Title: Structural organization of the human kininogen gene and a A;Reference number: A92545; WUID:85234583; PMID:2989294
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    100.0%; Score 84; DB 1; Length 427; 100.0%; Pred. No. 2.8e-06;
                                                                                                                                                                                                                                                               Fed. Proc. 27, 52-57, 1968
A,TLIE: Structural features of plasma kinins and kininogens. A,Reference number. A91455; MUID: 90255622; PMID:4952632
A,Contents: annotation; bradykinin
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                                                                                                                                                                                        A; Contents: annotation; gene organization
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C; Species: Rattus norvegicus (Norway rat)
C;Date: 20-Jun-1989 #sequence_revision 20-Jun-1989 #text_change 15-Nov-1996
C;Dates: 20-Jun-1989 #sequence_revision 20-Jun-1989 #text_change 15-Nov-1996
C;Accession: A28055
C;Accession: A28055
J; Bicl. Chem. 260, 12054-12059, 1985
A;Title: Primary structures of the mRNAs encoding the rat precursors for bradykinin a
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R; Kitagawa, H.; Kitamura, N.; Hayashida, H.; Miyata, T.; Nakanishi, S.
J. Biol. Chem. 262, 2190-2198, 1987
A; Tille: Differing expression patterns and evolution of the rat kininogen gene family A; Reference number: A92625; MUID:87137443; PMID:3029068
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C;Species: Rattus norvegicus (Norway rat)
C;Date: 08-Mar-1989 #sequence_revision 08-Mar-1989 #text_change 15-Nov-1996
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C;Species: Bos primigenius taurus (cattle)
C;Date: 14-Nov-1983 #sequence_revision 14-Nov-1983 #text_change 28-May-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      A Molecule type: mRNA
A; Residues: 1-639 <KIT>
A; Note: the authors translated the codon CAA for residue 347 as is
C; Superfamily: Kininogen; cystatin homology
C; Keywords: alternative splicing
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 433;
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A; Accession: A28055
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86.7%; Pred. No. 0.0029;
Mismatches 2;
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Pred. No. 0.002;
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A.Residues: 1-433 *FURA-
C.Superfamily: Kininogen; cystatin homology
C.Keywords: alternative splicing
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                                                                                                                                                                  K-kininogen, LMW I precursor - rat
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86.7%;
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Best Local Similarity
Matches 13; Conserv
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Matches 13; Conserv
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F.391-751-71-70-2000 May May Manual 1944 Chain #status experimental NEUL")
F.391-7510/Region: glycine/histidine/lysine-rich 30-residue repeats
F.421-510/Region: glycine/histidine/lysine-rich 30-residue repeats
F.431-44/Product: low molecular weight growth promoting factor #status experimental NET 197/Modified site: pyrrolidone carboxylic acid (Gln) (in mature form) #status experiments
F.197/Modified site: pyrrolidone carboxylic acid (Gln) (in mature form) #status experiments
F.197-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-380-70-38
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A; Residues: 264-359, 'N', 361-375 <LEN2>
A; Residues: 254-359, 'N', 361-375 <LEN2>
B; Little, S.S.; Johnson, D.A.
Biochem, J. 307, 341-346, 1995
A; Title: Human mast cell tryptase isoforms: separation and examination of substrate-specta-specta number: $55239; MUID:95251593; PMID:7733867
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A;Map position: 3427-3427
A;Nap position: 3427-3427
A;Nap position: 3427-3427
A;Nap position: 3427-3427
A;Nat position: 346/3; 375/3
C;Superfamily: Kininogen; cystatin homology
C;Superfamily: Kininogen; cystatin homology
C;Keywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; dupl.
F;19-18/Domain: signal sequence #status experimental <SIG>F;19-64/Product: HWW kininogen I (prokininogen) #status experimental <MATI>F;19-379,300-0014/Product: HWW kininogen II #status experimental <MATI>F;19-379/Domain: HWW kininogen heavy chain #status experimental <HCH>
F;19-131/Domain: cystatin homology <CYI>
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A; Residues: 431-434 <STR>
A; Residues: 431-434 <STR>
X; Kitamura, N:; Kitagawa, H.; Fukushima, D.; Takagaki, Y.; Miyata, T.; Nakanishi, S.
J. Biol. Chem. 260, 8610-8617, 1985
A; Title: Structural organization of the human kininogen gene and a model for its evoluti
A; Reference number: A92545; MUID:85234583; PMID:2989294
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Title: Purification from human plasma of a tetrapeptide that potentiates insulin-like A;Reference number: S68059; MUID:96033974; PMID:7589467
A;Accession: S68059
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F;264-375/Domain: cystatin homology <CY3>
F;380-389/Product: lysyl-bradykinin (kallidin II) #status experimental <KBDY>
F;381-389/Product: bradykinin (kallidin I) #status experimental <BDY>
R;Lenarcic, B.; Krasovec, M.; Ritonja, A.; Olafsson, I.; Turk, V. FEBS Lett. 280, 211-215, 1991
Affitle: Inactivation of human cystatin C and kininogen by human cathepsin D. A;Reference number: S14303; MUID:91192133; PMID:2013314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 A; Molecule type: protein
A;Residues: 450-452, X',454,'X',456 <LIT>
K:Straczek, J:; Maachl, F.; le Nguyen, D.; Becchi, M.; Heulin, M.H.; Nabet,
FEBS Lett. 373, 207-211, 1995
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Fed. proc. 27, 52-57, 1968
A:Title: Structural features of plasma kinins and kininogens. A:Reference number: A91455; MUID:90255622; PMID:4952632
A:Contents: annotation; bradykinin
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100.0%; Pred. No. 4.2e-06;
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Gaps

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C; Accession: A01284

293 NNATFYFKIDNVKKAR 308

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A Molecule type: protein

A Molecule type: p
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                                                                                                                                                                                 Hashimoto, N.; Kato, H.; Hayashida, H.; Miyata, T.; Iwana
                                        A.Accession: A91938
A;Molecule type: protein
A;Residues: 456-496 <HA2>
R;Suesolous: 456-496 <HA2>
J. Biol. Chem. 262, 2768-2779, 1987
A;Title: Bovine high molecular weight kininogen. The amino acid sequence, positions A;Reference number: A92627; MUD:87137530; PMID:3546295
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          major acute phase alpha-1 protein precursor - rat (fragment)
N;Contains: bradyfinin
C;Species: Rattus norvegicus (Norway rat)
C;Species: 27-Nov-1985 #sequence_revision 27-Nov-1985 #text_change 12-Apr-1996
C;Accession: A01285
R;Cole, T; Inglis, A.S.; Roxburgh, C.M.; Howlett, G.J.; Schreiber, G.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 59; DB 1; Length 619;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             2; Indels
             A; Reference number: A91938; MUID: 75170265; PMID: 1169237
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Pred. No. 0.061;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              :: ||||||| ||||
290 HDGTFYFKIDTVKKA 304
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Matches 11; Conserv
                                                                                                                                                                                                                                                                                                                                                 A; Accession: B29559
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
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                                                                                                                                                                                                   A; Molecule type: mRNA
A; Molecule type: mRNA
A; Molecule type: mRNA
A; Residues: 1-434 <NAMA
A; Cross-references : GB:V00427; GB:J00011; NID:g489; PIDN:CAA23710.1; PID:g490
A; Cross-references : GB:V00427; GB:J00011; NID:g489; PIDN:CAA23710.1; PID:g490
C; Comment: The LMW kininogen precursor is produced from the same gene as the HMW form as
C; Comment: Rininogen is a cysteine proteinase inhibitor, takes part in initiation of the
C; Comment: Rininogen is a cysteine protein prior to the release of bradykinin.
C; Superfamily: kininogen; cystatin homology
C; Reywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyd
C; Reywords: alternative splicing; blood coagulation; cysteine proteinase inhibitor; glyd
F; 19-337/Product: LMW kininogen II #status predicted <AICH>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                F.19-130/Domain: cystatin homology <CY1>
F.141-252/Domain: cystatin homology <CY2>
F.241-372-20main: cystatin homology <CY2>
F.241-372-20main: cystatin homology <CY2>
F.241-372-20main: cystatin homology <CY2>
F.377-386/Product: lysyl-bradykinin (kallidin II) #status predicted <BDY>
F.377-386/Product: LMW kinlinogen I light chain #status experimental <LCH>
F.387-434/Product: LMW kinlinogen I light chain #status experimental <LCH>
F.387-434/Product: LMW kinlinogen I light chain #status predicted complete stre: pyrcolidone carboxylic acid (Gln) (in mature form) #status predicted F.27-404,82-93,106-125,141-144,205-217,228-247,261-264,325-337,348-367/Disulfide bonds:
F.47,87.168.169.197,204,280/Blnding site: carbohydrate (Asn) (covalent) #status predicted F:380/Modified site: Met-Lys (kallikrein) #status predicted
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N'Contains: bradykinin (kallidin); kininogen I; kininogen II; prokininogen
C'Species: Bos primigenius taurus (cattle)
C'Date: 14-Nov-1983 #sequence_revision 14-Nov-1983 #text_change 22-Jun-1999
C'Accession: A01282; A91923; A91941; A91938 #22559
R'Kitamura, N.; Takagaki, Y.; Furuto, S.; Tanaka, T.; Nawa, H.; Nakanishi, S.
A'Ittle: A single gene for bovine high molecular weight and low molecular weight kininog
A'Reference number: A93317; MUID:84014106; PMID:6571699
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A; Residues: 387-455 <HAN>
A; Residues: 398-Pro, 401-Val, and 455-Lys were also found
B; Han, Y. N., Komiya, M.; Iwanaga, S.; Suzuki, T.
J. Biochem. 77, 55-68, 1975
A; Title: Studies on the primary structure of bovine high-molecular-weight kininogen. Amil
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A; Residues: 1-619 < KIT>
A; Cross-references: GB:V01492; GB:K01758; NID:g493; PIDN:CAA24736.1; PID:g494
A; Cross-references: GB:V01492; GB:K01758; NID:g493; PIDN:CAA24736.1; PID:g494
B; Kato, H.; Nagasawa, S.; Suzuki, T.
J. Biochem, 67, 313-323, 1970
A; Title: Studies on the structure of bovine kininogen: cleavages of disulfide bonds and A; Reference number: A91923; MUID:70180420; PMID:4986212
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                A; Molecule type: protein
A; Residues: 376-391 <KAT>
R; Han, TN. N; Kato, H; Iwanaga, S.; Suzuki, T.
Bilochem. 79, 1201-1222, 1976
A; Title: Primary structure of bovine plasma high-molecular-weight kininogen. The amino
A; Reference number: A91941; MUID:76260155; PMID:956151
R;Nawa, H.; Kitamura, N.; Hirose, T.; Asai, M.; Inayama, S.; Nakanishi, S. Proc. Natl. Acad. Sci. U.S.A. 80, 90-94, 1983 Astitle: Primary structures of bovine liver low molecular weight kininogen precursors A;Reference number: A93984; MUID:83117859; PMID:6572010
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ö
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 59; DB 1; Length 434;
Pred. No. 0.043;
2; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      kininogen, HMW II precursor - bovine
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73.3%;
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Matches 11; Conservative
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Gaps

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A;Cross-references: GB:M14356; NID:g205090; PIDN:AAA41492.1; PID:g205091
R;Enjyoji, K.; Kato, H.; Hayashi, I.; Oh-ishi, S.; Iwanaga, S.
J. Biol. Chem. 263, 965-972, 1988
A;Title: Purification and characterization of two kinds of low molecular weight kinin A;Reference number: A28525; MUID:88087225; PMID:333530
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ceding bradykinin.
C;Comment: T-kininogens contain T-kinin (I-S-bradykinin), a novel kinin isolated afte
d of an Arg or Lys, it is probably not released from its precursor by either tissue o
C;Comment: The T-kininogens are produced in response to an inflammatory stimulant.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                A) Introns: 65/3; 102/3; 130/1; 187/3; 223/2; 252/1; 309/3; 345/3; 374/3; 398/3

C; Superfamily: kininogen; cystatin homology

C; Superfamily: kininogen; cystatin homology

C; Superfamily: kininogen; pradykinin; cystatine proteinase inhibitor; duplication; glyc

F; 1-15/Domain: signal sequence #status predicted <SIG>

F; 19-430/Product: T-kininogen I #status experimental <MAT>

F; 19-130/Domain: cystatin homology <CXI>
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F:19Y/modified site: pyrrolidone carboxylic acid (Gln) (in mature form) #status experi
F:82,126,168,204,356/Binding site: carbohydrate (Asn) (covalent) #status predicted
F:83-94,107-125,141-144,205-217,228-247,263-266,327-339,350-369/Disulfide bonds: #sta
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             A:Experimental source: clone pSG17
C:Comment: At least three types of LMM kininogen precursors are present in rat plasma
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A.Residues: 5-430 <AND2>
A.Residues: 5-430 <AND2>
A.Cross-references: GB:M11661; NID:g205307; PIDN:AAA41570.1; PID:g205308
A.Note: the authors translated the codon CTC for residue 410 as Arg, CTA for residue C.Superfamily: kininogen; cystatin homology <CY1>
F:19-130/Domain: cystatin homology <CY2>
F:141-252/Domain: cystatin homology <CY2>
F:263-374/Domain: cystatin homology <CY3>
                                                                                                                                                                                                                                                                                                                                                                                                                                                   A; Molecule type: protein
A; Residues: 376-430 CBN2>
A; Residues: 376-430 CBN2>
A; Reinra, F.; Walter, R.; Vautravers, P.; Guigoz, Y.
Arch. Biochem. Biophys. 322, 333-338, 1995
A; Title: Identification of several isoforms of T-kininogen expressed in the liver of A; Reference number: S68034; MUD:96032652; PMID:7574705
A; Accession: S68036
A; Molecule type: mRNA
A; Residues: 340-430 <SIE>
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C;Species: Rattus norvegicus (Norway rat)
C;Date: 19-May-1989 #sequence_revision 19-May-1989 #text_change 20-Aug-1999
C;Date: 19-May-1989 #sequence_revision 19-May-1989 #text_change 20-Aug-1999
C;Accession: A23897; B23897
R;Anderson, K.P.; Heath, E.C.
J. Biol. Chem. 260, 12065-12071, 1985
A;Title: The relationship between rat major acute phase protein and the kininogens.
A;Reference number: A23897; MUID:86008266; PMID:2413019
        A; Reference number: A25488; MUID:87137465; PMID:3818598
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0.13;
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Pred. No. 0.13;
1; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         F;141-252/Domain: cystatin homology <CY2> F;263-374/Domain: cystatin homology <CY3>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        66.78;
73.38;
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Best Local Similarity 73.3
Matches 11; Conservative
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                                                                                                                          A; Molecule type: DNA
A; Residues: 1-48 <KAG>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A; Accession: B23897
                                                      A; Accession:
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R; Kageyama, R.; Kitamura, N.; Ohkubo, H.; Nakanishi, S.
J. Biol. Chem. 262, 2345-2351, 1987
A; Title: Differing utilization of homologous transcription initiation sites of rat K and
                                                                                                                                                        C;Superfamily: Kininogen; cystatin homology
C;Keywords: bradykinin; cysteine proteinase inhibitor; duplication; glycoprotein; inflam
C;Keywords: bradykinin; cysteine proteinase inhibitor; duplication; glycoprotein; inflam
F;1-11/Domain: signal sequence (fragment) *status predicted <MAT>
F;12-423/Product: major acute phase alpha-1 protein *status predicted <MAT>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                C; Species: Rattus norvegicus (Norway rat)
C; Date: 17-Mar-1987 #sequence_revision 17-Mar-1987 #text_change 22-Jun-1999
C; Date: 17-Mar-1987 #sequence_revision 17-Mar-1987 #text_change 22-Jun-1999
C; Accession: A01286; D25486; A28526; PL0193; JQ0027; B25488; A28525; S68036
R; Furuto-Kato, S.; Matsumoto, A.; Kiramura, N.; Nakanishi, S.
J. Blol. Chem. 260, 12054-12059, 1985
A; Title: Primary structures of the mRNAs encoding the rat precursors for bradykinin and
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A;Residues: 1:60,/E',62-113,'R',115-165,'F',167-178,'TKI',182-211,'F',213-256,'S',258-38
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (in mature form) #status predicted
, A;Molecule type: mRNA
A;Residues: 1-423 <COL>
C;Comment: This plasma glycoprotein inhibits cysteine proteinases. During acute inflamm
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A; Residues: 375-430 <KIT>
A; Residues: 375-430 <KIT>
B; Enjydji, K.; Kato, H.; Hayashi, I.; Oh-ishi, S.; Iwanaga, S.
J. Biol. Chem. 263, 973-979, 1988
A; Title: Purification and characterization of rat T-kininogens isolated from plasma of A; Reference number: A92729; MUID:88087226; PMID:3121623
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J. Biol. Chem. 262, 1910-2198
A;Title: Differing expression patterns and evolution of the rat kininogen gene family.
A;Reference number: A92625; WUID:87137443; PMID:3029068
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A; Residues: 'E', 20-48;376-430 <ENJ>
A; Residues: 'E', 20-48;376-430 <ENJ>
K; Kanda, S.; Sugiyama, K.; Takahashi, M.; Shumiya, S.; Tomino, S.; Nagase, S.
Jpn. J. Cancer Res. 81, 63-68, 1990
A; Title: Identification of a protein increasing in serum of Nagase analbuminemic rats
A; Reference number: PL0193; MUID:90216390; PMID:2108948
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A;Residues: 1-430 <FUR>
A;Cross-references: GB:M11883; NID:g205084; PIDN:AAA41489.1; PID:g205085
                                                                                                                                                                                                                                                                                                                    F;12-123/Domain: cystatin homology <CYi>
F;114-245/Domain: cystatin homology <CX2>
F;256-367/Domain: cystatin homology <CX3>
F;371-379/Product: bradykinin #status predicted <BDX>
F;371-379/Product: bradykinin #status predicted <BDX>
F;12/Modified site: pyrrolidone carboxylic acid (Gln) (in mature form) #F;12/Modified site: carbohydrate (Asn) (covalent) #status predicted
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 423;
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A;Residues: 330-420, R',422-429,'P' <KAN>
R;Anderson, K.P.; Croyle, M.L.; Lingrel, J.B.
Gene Bl, 119-128, 1989
A;Title: Primary structure of a gene encoding rat T-kininogen.
A;Reference number: JQ0027; MUID:90034172; PMID:2806908
A;Accession: JQ0027; MUID:90034172; PMID:2806908
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       A; Reference number: A92496; MUID:86008264; PMID:2413018
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Pred. No. 0.13;
1; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        N;Alternate names: 73K protein; LMW kininogen T-I
N;Contains: bradykinin; T-kinin
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73.38;
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Best Local Similarity
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Gaps ö

3; Indels

Length 430;

DB 2;

Score 56;

66.78;

Query Match

Length 430;

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Across references: GB:J00010; GB:V00426; NID:9163256; PIDN:AAA30604.1; PID:9163257 C; Comment: The LMW kininogen precursor is produced from the same gene as the HWW form C; Comment: Sininogen is a cysteine proteinase inhibitor, takes part in initiation of C; Comment: Bradykinin, released from kininogen by kallikrein, is a potent vasodilator xyproline residue is present in the kininogen prior to the release of bradykinin. C; Superfamily: kininogen; cystatin homology c; Superfamily: kininogen; cystatin homology c; Superfamily: kininogen; cystatin homology c; F:1-18/Domain: signal sequence #status predicted <SIG> F:19-436/Product: LMW kininogen I #status predicted <MAT> F:19-378/Product: LMW kininogen I heavy chain #status predicted <HCH>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Fig9-130/Domain: cystatin homology <CYI>
Fig9-130/Domain: cystatin homology <CYI>
Fig19-130/Domain: cystatin homology <CYI
Fig19-130/Domain: cystatin h
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Pred. No. 0.29;
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66.78;
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Best Local Similarity 66.7
Matches 10; Conservative
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292 HDGAFYFKIDTVKKA 306
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    C;Species: Rattus norvegicus (Norway rat)
C;Date: 20-Jun-1989 #sequence_revision 20-Jun-1989 #text_change 12-Dec-1997
C;Accession: B2805; E25486; B28526; C28526
R;Furuto-Kato, S.; Matsumoto, A.; Kiamura, N.; Nakanishi, S.
J. Biol. Chem. 260, 12054-12059, 1985
A;Title: Primary structures of the mRNAs encoding the rat precursors for bradykinin and
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A;Title: Primary structures of bovine liver low molecular weight kininogen precursors an A;Reference number: A93984; MUID:83117859; PMID:6572010
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A; Residues: 375-430 (KIT>
R:Enjyoji, K:, Kato, H:; Hayashi, I.; Oh-ishi, S.; Iwanaga, S.
J. Biol. Chem. 263, 973-979, 1988
A; Title: Purification and characterization of rat T-kininogens isolated from plasma, of
A; Reference number: A92729; MUID:88087226; PMID:3121623
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A; Residues: 1.430 < PCDR>
R; Kitagawa, H.; Kitamura, N.; Hayashida, H.; Miyata, T.; Nakanishi, S.
J. Biol. Chem. 262, 2190-2198, 1987
A; Title: Differing expression patterns and evolution of the rat kininogen gene family.
A; Reference number: A92625; MUID:87137443; PMID:3029068
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C; Date: 14-Nov.1983 #sequence_revision 14-Nov-1983 #text_change 22-Jun-1999
                                                       Gaps
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Pred. No. 0.13;
1; Mismatches 3; Indels
                                                   Indels
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A Residues: 'E'. 20-48 | 376-388,'R', 390-419,'ER', 422-430 < EN2>
C Superfamily: Kininogen: cystatin homology
C; Keywords: glycoprotein: pyroglutamic acid
F:1-18/Domain: signal sequence #status predicted <SIG>
F:19-430/Product: T-kininogen, LWM II #status experimental <A
F:19-130/Domain: cystatin homology <CXI>
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    Pred. No. 0.13;
                                                   Mismatches
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A; Residues: 'E', 20-25, 'MD', 28-48, 376-430 <ENJ>
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73.38;
73.38;
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                                              Conservative
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Best Local Similarity 73.3
Matches 11, Conservative
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HNHLFYFKIDTVKKA 306
                                                                                                                                      NNATEYFKIDNVKKA 15
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Best Local Similarity
Matches 11; Conserv
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Gaps

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Indels

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A; Accession: A01923
A; Molecule type: protein
A; Residues: 378-393 «KAT>
A; Residues: 378-393 «KAT>
B; Residues: 378-393 «KAT>
B; Residues: 378-393 «KAT>
A; Refidues: 378-393 «KAT>
A; Refidues: 458-498 «Har)
A; Reference number: A01938; MUID:75170265; PMID:1169237
A; Reference number: A01938
A; Residues: 458-498 «Har>
B; Residues: 458-498 «Har>
B; Sueyoshi, T.; Miyata, T.; Hashimoto, N.; Kato, H.; Hayashida, H.; Miyata, T.; Iwana
J. Biol. Chem. 262, 2768-2779, 1987
A; Ritle: Bovine high molecular weight kininogen. The amino acid sequence, positions of A; Reference number: A92627; MUID:87137530; PMID:3546295
                                                                                                                                                                                                                                                                                                                                                                                                                                                                A; Molecule type: mRNA
A; Residues: 1-621 <KIT>
A; Residues: 1-621 <KIT>
A; Cross-references: GB: V01491; GB: K01757; NID: 9491; PIDN: CAA24735.1; PID: 9492
R; Kato, H.; Nagasawa, S.; Suzuki, T.
J. Blochem. 67, 313-323, 1970
A; Title: Studies on the structure of bovine kininogen: cleavages of disulfide bonds a A; Reference number: A91923; MUID: 70180420; PMID: 4986212
                                                                                        C; Species: Bos primigenius taurus (cattle)
C; Date: 14-ROV-1983 * Requence_revision 14-Nov-1983 * text_change 22-Jun-1999
C; Accession: A01281; A91923; A91938; A92559
R; Kitamura, N.; Takagaki, Y.; Furuto, S.; Tanaka, T.; Nawa, H.; Nakanishi, S.
Nature 305, 545-549; 1983
A; Tille: A single gene for bovine high molecular weight and low molecular weight kini
A; Reference number: A93317; MUID: 84014106; PMID: 6571699
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Eur. J. Biochem. 152, 307-314, 1985
A:Title: The amino acid sequence of the light chain of human high-molecular-mass kini
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 A;Wolecule type: protein
A;Residues: 'Z'.20-133,'I',125-127,'I',129-378 <SUE>
R;Lottspeich, F.; Kellermann, J.; Henschen, A.; Foertsch, B.; Muller-Esterl, W.
Eur. J. Biochem. 152, 307-314, 1988
N;Alternate names: alpha-2-thiol proteinase inhibitor; preprokininogen N;Contains: bradykinin (kallidin); kininogen I; kininogen II; prokininogen
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33 NNLLFYSKFYNISKSR 48

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Cydcossion: AH1213
Rydlaser, P.; Frangeul, L.; Buchrieser, C.; Amend, A.; Baquero, F.; Berche, P.; Bloec
; Dominguez Bernal, G.; Duchaud, E.; Durand, L.; Dussurget, O.; Entian, K.D.; Fsihl,
D.; Jones, L.M.; Karst, U.
Science 294, 849-852, 2001
A;Authors: Kreft, J.; Kuhn, M.; Kunst, F.; Kurapkat, G.; Madueno, E.; Maitournam, A.;
A;Authors: Kreft, J.; Simoes, N.; Tierrez, A.; Vazquez-Boland, J.A.; Voss, H.; Wehla
A;Title: Comparative genomics of Listeria species.
A;Reference number: AB1077; MUID:21537279; PMID:11679669
A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-466 <GLA>
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             hypochetical protein At2937310 [imported] - Arabidopsis thaliana C; Species: Arabidopsis thaliana (mouse-ear cress) C; Species: Arabidopsis thaliana (mouse-ear cress) C; Date 02-Feb-2001 #sequence_revision 02-Feb-2001 #text_change 02-Mar-2001 C; Accession: B84791 R; Lin, X.; Kaul, S.; Rounsley, S.D.; Shea, T.P.; Benito, M.I.; Town, C.D.; Fujii, C.Y M.; Koo, H.; Wofffat, K. S.; Cronin, L.A.; Shen, M.; Won, M. S.E.; Umayam, L.; Tallon, euss, D.; Nierman, W.C.; White, O.; Eisen, J.A.; Salzberg, S.L.; Fraser, C.M.; Venter Nature 402, 761-768, 1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A;Title: Sequence and analysis of chromosome 2 of the plant Arabidopsis thaliana.
A;Reference number: A84420; WUID:20083487; PMID:10617197
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Cross-references: GB:NC_003210; PIDN:CAC99190.1; PID:q16410514; GSPDB:GN00177
                                                                                                                                                                                                                    lmol112 (imported) - Listeria monocytogenes (strain EGD-e)
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A;Cross-references: GB:AE002093; NID:g4056485; PIDN:AAC98051.1; GSPDB:GN00139
                                                                                                                                                                                                                TN916 ORF21 homolog lmo1112 [imported] - Listeria monocytogenes (strain EGD
C;Species: Listeria monocytogenes
C;Date: 27-Nov-2001 #sequence_revision 27-Nov-2001 #text_change 27-Nov-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     A;Map position: 2
C;Superfamily: Arabidopsis thallana hypothetical protein T8H10.30
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Pred. No. 45;
3; Mismatches
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C;Genetics:
A;Gene: lmo1112
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175 TYYTKCDNIESAR 187
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Best Local Similarity
7; Conserve
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Best Local Similarity
Matches 6; Conserv
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85 YYYKIDNIKQ 94
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A; Molecule type: DNA
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A: Reference number: A91153; MUID: 86030270; PMID: 40054110
A: Contents: annotation; Dovine cleavage sites; bovine carbohydrate binding sites
R: Sueyoshi, T.; Miyata, T.; Kato, H.: Iwanaga, S.
Seikagatu 56, 808, 1984
A: Title: Disulfide bonds in bovine HWM kininogens.
A: Contents: annotation; disulfide bonds
A: Reference number: A94300
A: Title: Disulfide bonds in bovine HWM kininogens.
A: Note: article in Japanese
C: Comment: The HWM kininogen by Ablibtor, takes part in initiation of the C: Comment: Mininogen is a cysteine proteinase inhibitor, takes part in initiation of the C: Comment: Mininogen is a cysteine proteinase inhibitor, takes part in initiation of the C: Comment: The Bytiche-Aistidiachylysine-rich region of HWM kininogen light chain is imported to a cysteine proteinase inhibitor; duplication of the C: Comment: The Bytiche-Aistidiachylysine-rich region of HWM kininogen light chain is imported to a cystein bomology of cystein predicted casts
C: Superfamily: Aininogen; cystatin homology cxiv.
E: Superfamily: Aininogen in the kininogen prior to the release of bradykinin.
C: Superfamily: Aininogen in the kininogen prior to the release of bradykinin.
C: Superfamily: Aininogen in the kininogen prior to the release of bradykinin.
C: Superfamily: Aininogen in the kininogen in the systems experimental cystatin homology cxiv.
F: 19-130/Domain: cystatin hom
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    Bacteriol. 183, 5709-5717, 2001
    Bacteriol. 183, Fr., Jaskunas, S., Zook, C.; Baltz, R.H.; Jaskunas, S.R.; Ayuthors: Yang, Y.; Young-Bellido, M.; Zhao, G.; Zook, C.; Baltz, R.H.; Jaskunas, S.R.; A;Ttle: Genome of the Bacterium Streptococcus pneumoniae Strain R6.
    A;Reference number: A97872; MUID:21429245; PMID:11544234

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R; Hoskins, J.A.; Alborn Jr., W.; Arnold, J.; Blaszczak, L.; Burgett, S.; DeHoff, B.S.; e, R.; LeBlanc, D.J.; Lee, L.N.; Lefkowitz, E.J.; Lu, J.; Matsushima, P.; McAhren, S.; y, P.; Sun, P.M.; Winkler, M.E.
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Pred. No. 0.42;
2; Mismatches 3; Indels
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Pred. No. 10;
2; Mismatches
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Best Local Similarity 50.0%;
Matches 8; Conservative
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292 HDGAFYFKIDTVKKA 306
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A;Molecule type: DNA
A;Residues: 1-217 <KUR>
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legionella
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MEDLINE-85122621; PubMed-6441591;
Ohkubo I., Kurachi K., Takasawa T., Shiokawa H., Sasaki M.;
"Isolation of a human cDNA for alpha 2-thiol proteinase inhibitor and its identity with low molecular weight kininogen.";
Biochemistry 23:5691-5697(1984).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        of
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Structural organization of the human kininogen gene and a model for
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                                                                                                                                                                                                                                                                                                                      Polo42; Polo43; 21-JUL-1986 (Rel. 01, Created) 01-FEB-1996 (Rel. 33, Last sequence update) 16-OCT-2001 (Rel. 40, Last annotation update) Rininogen precursor (Alpha-2-thiol proteinase inhibitor) (Contains: Bradykinin).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Takagaki Y., Kitamura N., Nakanishi S.; "Cloning and sequence analysis of cDNAs for human high molecular weight and low molecular weight prekininogens. Primary structures
                                                                                                                                                                                                                                                                                                                                                                                                                                                     Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Kitamura N., Kitagawa H., Fukushima D., Takagaki Y., Miyata T.,
Nakanishi S.;
P45477 P31765 P31765 P31765 P67531 P40902 P67531 P40902 P67531 P7511 P75
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MEDLINE-86030270; PubMed-4054110;
Lottspeich F., Kellermann J., Henschen A., Foertsch B.,
Mueller-Esterl W.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Pierce J.V.; "Structural features of plasma kinins and kininogens.";
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        SEQUENCE FROM N.A. (ISOFORMS HMW AND LAWW).
                                                                                                                                                                                                                                   ALIGNMENTS
                                                MLTA_BUCAI
ISP7_SCHPO
Y16R_MYCIO
                                                                                               PROA_LEGLO
TFC1_YEAST
EFG_MICLU
PMA2_LYCES
                 GALM_HAEIN
VP26_CAEEL
                                                                                                                                                                   CC16_YEAST
                                                                                                                                                                                  TERT_OXYTR
                                                                                                                                                                                                                                                                                                                   PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      two human prekininogens.";
J. Biol. Chem. 260:8601-8609(1985).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            its evolution.";
J. Biol. Chem. 260:8610-8617(1985).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            mass kininogen.";
Eur. J. Biochem. 152:307-314(1985)
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MEDLINE-85234582; PubMed=2989293;
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                                                                                                                                                                                                                                                                                                                   STANDARD;
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SEQUENCE OF 381-389.
NCBI_TaxID=9606;
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 RESULT 1
KNG_HUMAN
rattus norv
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bos taurus
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Copyright (c) 1993 - 2003 Compugen Ltd.
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KININOGEN HEAVY CHAIN.

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RL SGİKAGAKU 56:808-808(1984).

C. !- FUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)

THW-KININOGEN PLAYS AN IMPORTANT ROLE IN BLOOD COAGULATION BY
THE HWA-KININOGEN INHIBITS THE THROMBIN-AND PLASMIN-
C. FACTOR XII; (3) HWM-KININOGEN INHIBITS THE THROMBIN-AND PLASMIN-
C. FACTOR XII; (3) HWM-KININOGEN INHIBITS THE THROMBIN-AND PLASMIN-
C. FACTOR XII; (4) HWM-KININOGEN INHIBITS THE THROMBIN-AND PLASMIN-
C. FACTOR XII; (5) HWM-KININOGEN INHIBITS THE ACTIVE PEPTIDE
C. FACTOR AGRECATION OF THROMBOCYTES; (4) THE ACTIVE PEPTIDE
C. PHYSIOLOGICAL EFFECTS: (4) INFLUENCE IN SHOWS A VARIETY OF
PHYSIOLOGICAL EFFECTS: (4) INFLUENCE IN SHOWN (4C) NATRIURESIS AND
C. CONTRACTION, (4B) INDUCTION OF HYPOTENSION, (4C) NATRIURESIS AND
C. DIURESIS, (4D) DECREASE IN BLOOD GLUCOSE LEVEL, (4E) IT IS A
MEDIATOR OF INFLAMMATION OF NOCICEPTORS (4E3) RELEASE OF
C. CHHER MEDIATORS OF INFLAMMATION (E.G. PROSTAGLANDINS), (4F) IT HAS
C. CONTRACTLY VIA ENDOTHELIUM PERIVED FRLAXING FACTION); (5)
LMW-KININOGEN INHIBITS THE AGGREGATION OF THROMBOCYTES; (6) LMW-KININOGEN IS IN CONTRAST TO HWM-KININOGEN NOT INVOLVED IN BLOOD
                                                                                                                                                                                                                                                                                                                                      This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation-the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@ib-sib.ch).
                                                                                                                                                                                                                                            -!- SUBCELLULAR LOCATION: Secreted.
-!- ALTERNATIVE PRODUCTS: 2 ISOFORMS; HWW (SHOWN HERE) AND LAWW; ARE PRODUCED BY ALTERNATIVE SPLICING.
-!- TISSUE SPECIFICITY: PLASWA.
-!- PIM: BRADYKININ IS RELEASED FROM KININGEN BY PLASMA KALLIKREIN.
-!- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Glycoprotein; Plasma; Repeat; Thiol protease inhibitor; Vasodilator; Bradykinin; Blood coagulation; Inflammatory response; Signal; Alternative splicing.
SIGNAL
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InterPro; IPR002395; Kininogen.
Pfam; PF00031; cystatin; 3.
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M11522; AAB59550.1;
M11523; AAB59550.1;
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EMBL; M11438; AAB59551.1;
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EMBL; M11522; AAB59551.1;
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M11525; AAB59551.1;
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M11527; AAB59551.1;
M11528; AAB59551.1;
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EMBL; M11438; AAB59550.1;
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M11525; AAB59550.1
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PIR; A02576; A25276.
PIR; A01280; KGHUL1.
PIR; B25276; B25276.
PIR; S02482; S02482.
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EMBL; M11528;
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EMBL; M11528;
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N-LINKED (GLCNAC. ..) (POTENTIAL).
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PPKAGARPASRREVS (IN ISOFORM LAW).
MISSING (IN ISOFORM LAW).
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Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
NCBL_TaxID=10116;
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BRADYKININ.
KININGEN LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
HIS-RICH (ASSOCIATED WITH CLOTTING ACTIVITY).
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MEDLINE-87137443; PubMed=3029068;
Kitagawa H., Kitamura N., Hayashida H., Miyata T., Nakanishi S.;
"Differing expression patterns and evolution of the rat kininogen
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ö
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100.0%; Pred. No. 1.3e-06;
....matches 0; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       -> I (IN REF. 1).
3132B4CBAF8FBB7E CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                  INTERCHAIN
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Best Local Similarity 100.
Matches 16; Conservative
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644 AA;
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KININOGEN

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Rageyama R., Kitamura W., Ohkudo H., Nakanishi S.;

"Differing utilization of homologous transcription initiation sites
"Differing utilization of homologous transcription initiation sites
"L. at K and T kininogen genes under inflammation condition.";
"J. Biol. Chem. 262:2345-2331(1987).

"I BIOL. Chem. 262:2345-2331(1987).

"I BIOL. Chem. 262:2345-2331(1987).

"I WAY KININOGEN PLAYS AN IMPORTANT ROLE IN BLOOD COAGULATION BY
HELPING TO POSITION OPTHALLY PREAALLIKREIN AND FACTOR XI NEXT TO
HELPING TO POSITION OPTHALLY PREAALLIKE THE THROUGH NEW A VARIETY OF
HEADYKININ THAT IS RELEASED FROM HUM-KININOGEN SHOWS A VARIETY OF
HYSTOCOGICAL EFFECTS: (4A) INFLIENCE IN SMOOTH MUSCLE
CONTRACTION, (4B) INDUCTION OF HYPOTENSION, (4C) NATRINESIS AND
DUNRESIS, (4D) DECREASE IN BLOOD GLUCOSE LEWEL, (4E) IT IS A
MEDIATOR OF INFLAMMATION OF NOCIEPTORS (4E) IT IS A
MEDIATOR OF INFLAMMATION OF OCCEPTORS (4E) IT HAS
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COTHER MEDIATORS OF INFLAMMATION OF OCCEPTORS (4E) IN HAS
COTHER MEDIATORS OF INFLAMMATION OF OCCEPTORS (4E) IN HAS
COTHER MEDIATORS OF INFLAMMATION OF THOMOGEN OF THROUGH IN BLOOD
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   TISSUE SPECIFICITY: PLASMA.

THE BRADAVENINI IS RELEASED FROM KININOGEN BY PLASMA KALLIKREIN.

MISCELLANGUS: RAT EXPRESS FOUR TYPES OF KININOGENS: THE CLASSICAL

HMM/LMM KININOGENS AND TWO ADDITIONAL LMW-LIKE KININOGENS: TI AND
  for major acute phase alpha 1-
   Glycoprotein; Plasma; Repeat; Thiol protease inhibitor; Vasodilator; Bradykinin; Blood coagulation; Inflammatory response; Signal; Alternative spliting; Multigene family.
   SUBCELLUIAR LOCATION: Secreted.
ALTERNATIVE PRODUCTS: 2 ISOFORMS; HWW (SHOWN HERE) AND LAW; ARE
PRODUCED BY ALTERNATIVE SPLICING.
"Primary structures of the mRNAs encoding the rat precursors for bradykinin and T-kinin. Structural relationship of kininogens with major acute phase protein and alpha 1-cysteine proteinase
  SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
  Fung W.-P., Schreiber G.;
"Structure and expression of the genes for major protein (thiostatin) and kiningen in the rat.";
J. Biol. Chem. 262:9298-9308(1987).
   EMBL; M14369; AAA41484.1; -
EMBL; M14369; AAA41485.1; -
EMBL; M1455; AAA41482.1; -
FIR; A22486; A25486.
PIR; A28055; A28055;
Interpro: IPROMOMOLO; Cystatin.
Interpro: IPROMOMOLO; Cystatin.
Interpro: IPROMOMOLO; Cystatin.
PFMR; PROMOMA; Cystatin; 3.
PRINTS; PROMOMA; CY; 3.
PROSITE; PSO0287; CY: 3.
  Biol. Chem. 260:12054-12059(1985)
  SEQUENCE OF 1-41 FROM N.A.
STRAIN-Wistar; TISSUE-Liver;
MEDLINE-87137465; PubMed-3818598;
   STRAIN-Buffalo;
MEDLINE-87250580; Pubmed-2439509;
   EMBL; L29428; AAA41486.1; -.
EMBL; M11884; AAA41487.1; -.
   SEQUENCE OF 1-65 FROM N.A.
  CLOTTING
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ö
   INTERCHALM (BY SIMILARITY).

BY SIMILARITY.

B
  MEDLINE-87137530; PubMed-3546295;
MEDLINE-87137530; PubMed-3546295;
MEDLINE-87137530; PubMed-3546295;
Miyata T., Hashimoto N., Kato H., Hayashida H.,
Miyata T., Lwanaga S.;
"Bovine high molecular weight kininogen. The amino acid sequence,
positions of carbohydrate chains and disulfide bridges in the heavy
chain portion.";
J. Biol. Chem. 262:2768-2779(1987).
-I-FUNCTION: (1) KININGENS ARE INHIBITORS OF THIOL PROTEASES; (2)
LIMW-KININOGEN INHIBITS THE AGGREGATION OF THROMBOCYTES; (3) THE
ACTIVE PEPTIDE KALLIDIN THAT IS RELEASED FROM LAW-KININOGEN SHOWS
A VARIETY OF PHYSIOLOGICAL EFFECTS: (3A) INFLUENCE IN SMOOTH
  Gaps
  MEDLINE-8117859; PubMed-6572010; MEDLINE-8117859; PubMed-6572010; Nawa H., Kitamura N., Hirose T., Asai M., Inayama S., Nakanishi S.; Nayamary structures of bovine liver low molecular weight kininogen precursors and their two mRNAs."; Proc. Natl. Acad. Sci. U.S.A. 80:90-94(1983).
  Bos taurus (Bovine).
Sukaryota: Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
Bovidae; Bovinae; Bos.
  ö
   Length 639;
  2; Indels
  MISSING (IN ISOFORM LMW).
E -> K (IN REF. 2).
D3172DF94FF56AF5 CRC64;
                  KININOGEN HEAVY CHAIN.
   KININOGEN LIGHT CHAIN.
   Score 67; DB 1; L
Pred. No. 0.00095;
  CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
HIS-RICH.
  Mismatches
   ö
  434 639 N
61 61 E
639 AA; 70933 MW;
   79.8%;
86.7%;
  Conservative
  293 NNHTEYEKIDTVKKA 307
  1 NNATFYFKIDNVKKA 15
  Similarity
  NCBI_TaxID-9913;
   Query Match
Best Local Simi
Matches 13;
   Bradykinin]
  DISULFID
  DISULFID
  CARBOHYD
VARSPLIC
  VARSPLIC
   DISULFID
   DISULFID
  DISULFID
  DISULFID
   DISULFID
   CARBOHYD
   CARBOHYD
  CARBOHYD
   CARBOHYD
   CARBOHYD
   SEQUENCE
                     CHAIN
PEPTIDE
   DOMAIN
   DOMAIN
   DOMAIN
   CHAIN
  RESULT 3
SO SELLE  셤
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   ö
MUSCLE CONTRACTION, (3B) INDUCTION OF HYPOTENSION, (3C)
NATRIURESIS AND DIURESIS (KIDNEY).
SUBCELLULAR LOCATION: Extracellular.
ALTERNATUP PRODUCTS: 2 isoforms; LMW II (shown here) and HMW II
(AC POLO45); are produced by alternative splicing.
TISSUE SPECIFICITY: PLASMA.
   Gaps
  -!- PTM: BRADYKININ IS RELEASED FROM KININOGEN BY PLASMA KALLIKKEIN.
-!- MISCELLANEOUS: LAM-KININOGEN IS IN CONTRAST TO HMW-KININOGEN NOT
   21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
Kininogen, HWW II precursor (Thiol proteinase inhibitor) (Contains: Bos taurus (Bovine)
  InterPro; IPR000010; Cystatin.
Fram; PR00031; Cystatin; 3.
SMART; SM00043; CY; 3.
PROSITE; PS00287; CYSTATIN; 2.
BROSITE; PS00287; CYSTATIN; 2.
Alycoprotein; Plasma; Repeat; Vasodilator; Alternative splicing; Thiol protease inhibitor; Bradykinin; Signal.
  CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
PYRROLIDONE CARBOXYLIC ACID.
N-LINKED (GLCNAC. . .).
O-LINKED (GLCNAC. . .); OR 169.
N-LINKED (GLCNAC. . .); OR 169.
N-LINKED (GLCNAC. . .); OR 169.
   ö
  Score 59; DB 1; Length 434; Pred. No. 0.014;
  2; Indels
  48148 MW; 73A7079DE3E03430 CRC64;
  N-LINKED (GLCNAC. . .).
N-LINKED (GLCNAC. . .).
  INVOLVED IN BLOOD CLOTTING.
-!- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
   KININGEN, LMW II. HEAVY CHAIN.
  619 AA.
  2; Mismatches
  LIGHT CHAIN
  INTERCHAIN
   BRADYKININ
  PRT;
   EMBL; V00427; CAA23710.1; -. PIR; A01284; KGBOL2. HSSP; P01038; 1A90.
  70.28;
73.38;
  Conservative
   STANDARD;
  1 NNATEYFKIDNVKKA 15
  434 AA;
  Best Local Similarity
   KNH2_BOVIN
P01045;
21-JUL-1986 (
21-JUL-1986 (
15-JUN-2002 (
   CHAIN
DOMAIN
DOMAIN
MOD_RES
CARBOHYD
  DISULFID
  CHAIN
PEPTIDE
   CARBOHYD
   CARBOHYD
  CARBOHYD
   CARBOHYD
  DISULFID
   DISULFID
  DISULFID
  DISULFID
  SULFID
  DISULFID
   DISULFID
  SEQUENCE
   Query Match
  CARBOHYI
   RESULT 4
KNH2_BOVIN
   CHAIN
  Matches
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  염
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   REAL REDUILE "DEVOLUE); FURMENCE-97011;

RET Han Y Y.N., Kato H., Iwanaga S., Suzuki T.;

RET Perimary structure of bovine plasma high-molecular-weight kininogen.

RET following the C-terminus of the bradykinin moiety.";

Following the C-terminus of the bradykinin moiety.";

Following the C-terminus of the bradykinin moiety.";

RET Following the C-terminus of the bradykinin moiety.";

RET Following the C-terminus of the bradykinin moiety.";

RET Following the C-terminus of the bradykinin moiety.";

RET Following the C-terminus of the bradykinin moiety.";

RET Following the C-terminus of the bradykinin moiety.";

RET Following to the primary structure of bovine high-molecular-weight kininogen. Amino acid sequence of a fragment ('histidine-rich RT kininogen. Amino acid sequence of a fragment ('histidine-rich RT kininogen. Amino acid sequence of a fragment ('histidine-rich RT kininogen. Amino acid sequence of a fragment ('histidine-rich RT Kininogen. Amino acid sequence of a fragment ('histidine-rich HT TRININOGEN PLAYS AN IMPORENT REDUIT OF CC HELPING TO POSITION OF THROMENT REDUIT OF CC HELPING TO POSITION OF THROMENT REDUIT OF SHOWS AND PLANINININOUS AND PLANINININOUS AND PLANINININOUS AND CAUSES (4) INPLOENCE IN SMOOTH MUSTICE CONTRACTION, (4B) INDUCTION OF HYDOTHORIC IN SMOOTH MUSTICE TO PACTOR MISCISSIS AND DIURESIS, (4B) INDUCTION OF NOCIDEPTORS (4E) IN TARANDENT OF NOCIDEPTORS (4F) IN TARA A CARDIOPROTER EFFECT (DIRECTLY VIA BRADIATINE ACTION, INDIRECTLY VIA ENDOTHELIUM-DERIVED RELAXING FACTOR.

ANTONIN INDIRECTLY VIA ENDOTHELIUM-DERIVED RELAXING FACTOR.
   L1.)
SEQUENCE FROM N.A.
MEDLIND=84014106; PubMed=6571699;
Kitamura N., Takagaki Y., Furuto S., Tanaka T., Nawa H., Nakanishi S.;
"A single gene for bovine high molecular weight and low molecular
   MEDLINE-70180420; PubMed-4986212;
Kato H., Nagasawa S., Suzuki T.;
"Studies on the structure of bovine kininogen: cleavages of disulfide
bonds and of methionyl bonds in kininogen-II.";
J. Biochem. 67:313-323(1970).
   (AC POLO47); are produced by alternative splicing.
--- TISSUE SPECIFICITY: PLASMA.
--- PTH: BRADYKININ IS RELEASED FROM KININGGEN BY PLASMA KALLIKREIN.
--- SIMILARITY: CONTAINS 3 CYSTAIN-LIKE DOMAINS.
  "Bovine high molecular weight kininogen. The amino acid sequence, positions of carbohydrate chains and disulfide bridges in the heavy
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea; Bovidae; Bovinae; Bos.
  SEQUENCE OF 19-376.
MEDILINE-87137530; PubMed-3546295;
SUEYOSHI T., Miyata T., Hashimoto N., Kato H., Hayashida H.,
Miyata T., Iwanaga S.;
   SUBCELLULAR LOCATION: Extracellular.
  J. Biol. Chem. 262:2768-2779(1987).
   MEDLINE-76260155; PubMed-956151;
   SEQUENCE OF 376-391.
   SEQUENCE OF 387-455.
   NCBI_TaxID=9913
  chain portion.
   ACTION)
```

```
major acute phase protein and alpha 1-cysteine proteinase
  KALLIKREIN
   PROTEASES
   kininogens.
 ö
   Eukaryota; Métazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalla; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
  Gaps
  SEQUENCE FROM N.A.

SEQUENCE FROM N.A.

FUTUTO-KATO S., MATSUMOTO A., Kitamura N., Nakanishi S.;

FUTUTO-KATO S., MATSUMOTO A., Kitamura N., Nakanishi S.;

"Pilmary structures of the mRNAs encoding the rat precursors for bradykinin and T-kinin. Structural relationship of kininogens with
   Glycoprotein; Plasma; Repeat; Vasodilator; Alternative splicing; Thiol protease inhibitor; Bradykinin; Blood coagulation; Signal; Inflammatory response.
  ö
  (PARTIAL).
  PTROLLIDONE CARBOXYLIC ACID.
N-LINKED (GLCNAC. . ).
N-LINKED (PARTIAL).
N-LINKED (GLCNAC. . ); OR 169.
N-LINKED (GLCNAC. . ) (PARTIAL).
N-LINKED (GLCNAC. . ).
   70.2%; Score 59; DB 1; Length 619;
   2; Indels
  998 T -> P.
101 L -> V.
154 H -> K.
68710 MW; F04320A8EBOEEODA CRC64;
   KININOGEN, HMW II
HEAVY CHAIN.
  LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
  Pred. No. 0.02; ; Mismatches
   430 AA.
   BRADYKININ.
   INTERCHAIN.
  O-LINKED
   PRT;
EMBL; V01492; CAA24736.1; -.
EMBL; V01492; CAA24737.1; ALT_SEQ.
   5;
                           HSSP; P01038; 1A90.
InterPro; IPR000010; Cystatin.
InterPro; IPR002395; Kininogen.
Pfam; PF00031; cystatin; 3.
PRINTS; PR00334; KININOGEN.
  PROSITE; PS00287; CYSTATIN; 2.
   Conservative
   STANDARD;
  :: ||||||| ||||
290 HDGTFYFKIDTVKKA 304
   Rattus norvegicus (Rat).
  1 NNATFYFKIDNVKKA 15
  93
   200
280
580
589
              PIR; A01282; KGBOH2.
PIR; B29559; B29559.
  SM00043; CY;
   27
82
106
141
205
228
228
325
348
398
454
619 AA;
   Best_Local Similarity
Matches 11; Conserv
  NCBI_TaxID=10116;
   MOD_RES
CARBOHYD
  DISULFID
DISULFID
  CHAIN
CHAIN
PEPTIDE
  CARBOHYD
  CARBOHYD
   CARBOHYD
   CARBOHYD
  CARBOHYD
  CARBOHYD
   DISULFID
   DISULFID
  DISULFID
   DISULFID
  DISULFID
   DISULFID
   DISULFID
   SEQUENCE
   Query Match
   KNT1_RAT
  VARIANT
VARIANT
VARIANT
   CHAIN
  SMART;
  SIGNAL
  DOMAIN
  DOMAIN
g
   ð
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  -I- MISCELLANEOUS: RAT EXPRESS FOUR TYPES OF KININOGENS: THE CLASSICAL HISCELLANEOUS: RAT EXPRESS FOUR TYPES OF KININOGENS: THE CLASSICAL HWA AND LAW KININOGENS THE SAME GENE, AND TWO ADDITIONAL LAW-LIKE KININOGENS: T-I AND T-II.
-I- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
-I- CAUTION: IN ADDITION TO THE CONFLICTS DESCRIBED IN THE FEATURE TABLE, REF. 2. SEQUENCE DIFFERS FROW THAT SHOWN IN POSITIONS: 257, 262, 268, 269, 295, 314, 312, 314, 332, AND 389. IN ALL THOSE POSITIONS THE ALTERNATE AMINO-ACID IS THE ONE PRESENT IN T-II KININOGEN.
   SEQUENCE OF 7-430 FROM N.A.

MEDLINE-85127561; Pubmed=2578992;

Cole T., Inglis A.S., Roxburgh C.M., Howlett G.J., Schreiber G.;

"Major acute phase alpha 1-protein of the rat is homologous to bovine kininopen and contains the sequence for bradykinin: its synthesis is regulated at the mRNA level.";

FBBS Lett. 182:57-61(1985).
  T-KININOGEN I IS RAISED.
-!- PTM: AS T-KININ IS PRECEDED BY A MET INSTEAD OF AN ARG OR LYS, IT IS NOT RELEASED FROM ITS PRECURSOR BY EITHER TISSUE OR PLASMA
   Fung W.-P., Schreiber G.; "Structure and expression of the genes for major acute phase alpha 1-
   -i-TISSUE SPECIFICITY: PLASMA.
-i-INDUCTION: IN RESPONSE TO AN INFLAMMATORY STIMULANT. T-KININGEN
II SYNTHESIS IS INDUCED AND THE PLASMA CONCENTRATION OF
  Anderson K.P., Heath E.C.; "The relationship between rat major acute phase protein and the
  PROSITE; PS00287; CYSTATIN; 2.
Glycoprotein; Plasma; Repeat; Vasodilator; Multigene family;
Thiol protease inhibitor; Bradykinin; Acute phase; Signal.
1 16
   SEQUENCE OF 5-430 FROM N.A., AND PARTIAL SEQUENCE
   -1- SUBCELLULAR LOCATION: Extracellular.
  Biol. Chem. 260:12065-12071(1985).
Biol. Chem. 260:12054-12059(1985)
  EMBL; X02299; CAA26162.1; ALT_SEQ.
  MEDLINE-86008266; PubMed-2413019;
   SEQUENCE OF 1-65 FROM N.A. MEDLINE-87250580; PubMed-2439509;
  GlycoSuiteDB; P01048; -.
InterPro; IPR000010; Cystatin.
Pfam: PF00031; cystatin; 3.
SMART; SM00043; CY; 3.
  EMBL; M11883; AAA41489.1; -.
EMBL; M11661; AAA41570.1; -.
EMBL; M16454; AAA41568.1; -.
  PIR; A01286; KGRTT1
PIR; A01285; KGRTM.
   PIR; A23897; A23897
PIR; A27115; A27115
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   entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
   -i- MISCELLANEOUS: RAT EXPRESS FOUR TYPES OF KININOGENS: THE CLASSICAL HAW AND LAW KININOGENS PRODUCED BY ALTERNATIVE SPLICING OF THE SAME GENE, AND TWO ADDITONAL LAW-LIKE KININOGENS: T-I AND T-II.
-i- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
                    T-KININOGEN I IS RAISED.
-!- PTM: AS T-KININ IS PRECEDED BY A MET INSTEAD OF AN ARG OR LYS, IT IS NOT RELEASED FROM ITS PRECURSOR BY EITHER TISSUE OR PLASMA
  -i-TISSUE SPECIFICITY: PLASMA.
-i-INDUCTION: IN RESPONSE TO AN INFLAMMATORY STIMULANT. T-KININGEN
II SYNTHESIS IS INDUCED AND THE PLASMA CONCENTRATION OF
  bradykinin and T-kinin. Structural relationship of kininogens with
   INTERCHAIN (BY SIMILARITY).

BY SIMILARITY.

N-LIKED (GLCNAC. ...) (POTENTIAL).

N-LINKED (GLCNAC. ...) (POTENTIAL).
  (POTENTIAL)
  ö
  PROSITE; PS00287; CYSTATIN; 2.
Glycoprotein; Plasma; Repeat; Vasodilator; Multigene family;
Thiol protease inhibitor; Bradykinin; Acute phase; Signal.
1 18
            major acute phase protein and alpha 1-cysteine proteinase
   Length 430;
  3; Indels
   43EDF02D1BF55076 CRC64;
   LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
   Score 56; DB 1;
Pred. No. 0.045;
  KININOGEN, T-II.
  (GLCNAC
  Mismatches
  HEAVY CHAIN.
  SUBCELLULAR LOCATION: Extracellular.
   LINKED
  T-KININ-T
  EMBL; M11885; AAA41491.1; -.
PIR; B28055; B28055.
G1ycoSuiteDB; P08932; -.
InterPro; IPR000010; Cystatin.
Pfam; PF00031; cystatin; 3.
SMART; SM00043; CY; 3.
   47524 MW;
  Conservative
   : | |||||| |||| |||| |||| : 292 HNHLFYFKIDTVKKA 306
   1 NNATFYFKIDNVKKA 15
   375
375
3386
430
135
2257
2257
404
125
125
217
   369
82
126
168
204
326
  141
205
203
328
327
350
82
126
168
326
430 AA;
   Query Match
Best Local Similarity
   KALLIKREIN
   11;
  DOMAIN
DOMAIN
DOMAIN
DISULFID
   DISULFID
DISULFID
CARBOHYD
   DISULFID
DISULFID
DISULFID
   DISULFID
DISULFID
  CARBOHYD
  CARBOHYD
SEQUENCE
  CHAIN
CHAIN
PEPTIDE
CHAIN
   DISULFID
  CARBOHYD
   Matches
  q
   á
   ö
   Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
   Gaps
  01-NOV-1988 (Rel. 09, Created)
01-NOV-1988 (Rel. 09, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
T-kininogen II precursor (Major acute phase protein) (Alpha-1-MAP)
(Thiostatin) [Contains: T-kinin].
   (POTENTIAL).
(POTENTIAL).
(POTENTIAL).
(POTENTIAL).
  SEQUENCE FROM N.A.
MEDILINE-886008204; Fulbwed-2413018;
Furuto-Kato S., Matsumoto A., Kitamura N., Nakanishi S.;
"Primary structures of the mRNAs encoding the rat precursors for
   (POTENTIAL)
  ó
  INTERCHAIN.

BY SIMILARITY.
   Score 56; DB 1; Length 430;
Pred. No. 0.045;
.; Mismatches 3; Indels
   NC - MDR (IN REF. 2).

V -> L (IN REF. 2).

C -> Y (IN REF. 3).

C -> Y (IN REF. 3).

S -> F (IN REF. 2).

REV -> TRI (IN REF. 2).

R -> F (IN REF. 2).

S -> F (IN REF. 2).

S -> F (IN REF. 2).

S -> F (IN REF. 2).

T -> R (IN REF. 2).

R -> F (IN REF. 2).

R -> TM (IN REF. 2).

S -> TM (IN REF. 2).
  -> Q (IN REF. 2).
-> G (IN REF. 2 AND 3).
-> L (IN REF. 2).
H -> ER (IN REF. 3).
   FAEBB78FAF4723C3 CRC64;
  CYSTATIN-LIKE 1. CYSTATIN-LIKE 2. CYSTATIN-LIKE 3.
                                    CHAIN.
   DH -> ER
  Ŷ
  47715 MW;
  66.7%;
73.3%;
  Local Similarity 73.3
  292 HNHIFYFKIDTVKKA 306
  STANDARD;
   1 NNATFYFKIDNVKKA 15
  82
126
168
168
326
28
28
55
61
166
166
193
  430 AA;
   NCBI_TaxID-10116;
  DISULFID
   DISULFID
  DISULFID
  CARBOHYD
CARBOHYD
  CONFLICT
CONFLICT
CONFLICT
CONFLICT
CONFLICT
  CONFLICT
  CONFLICT
  KNT2_RAT
  DISULFID
  DISULFID
  DISULFID
   CARBOHYD
   CARBOHYD
   CARBOHYD
  SEQUENCE
   Query Match
   CONFLICT
   CONFLICT
                         PEPTIDE
                                 CHAIN
   DOMAIN
DOMAIN
  Matches
   KNT2_RAT
  RESULT
ò
  a
```

Gaps

```
KNH1_BOVIN
   DISULFID
  CONFLICT
                      CARBOHYD
                                     DISULFID
   DISULFID
  DISULFID
   DISULFID
  DISULFID
   DISULFID
   DISULFID
  P01044;
   KNH1_BOVIN
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   JUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)
-1- FUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)
-1- FUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)
-1- KININOGEN INHIBITS THE AGENEGATION OF THROMBOCYTES; (3) THE
ACTIVE PEPTIDE KALLIDIN THAT IS RELEASED FROM LAW-KININOGEN SHOWS
A VARIETY OF PHYSIOLOGICAL EFFECTS: (3A) INFLUENCE IN SMOOTH
MUSCLE CONTRACTION, (3B) INDUCTION OF HYPOTENSION, (3C)
-1- SUBCELLUAR LOCATION: Extracellular.
-1- SUBCELLUAR LOCATION: Extracellular.
-1- ALTERNATIVE PRODUCTS: 2 isoforms; LAW I (shown here) and HAW I
-1- TISSUE SPECIFICITY: PLASAM.
-1- TISSUE SPECIFICITY: PLASAM.
-1- TISSUE SPECIFICITY: PLASAM.
-1- PIW: BRADYKININ IS RELEASED FROM KININOGEN BY PLASAM. KALLIKREIN.
-1- MISCELLANBOUS: LAW-KININOGEN IS IN CONTRAST TO HAW-KININOGEN NOT
   SEQUENCE OF 19-378.
MEDLINE-81137530; PubMed-3546295;
Sueyoshi T., Miyata T., Hashimoto N., Kato H., Hayashida H.,
Miyata T., Iwanaga S.;
"Bovine high molecular weight kininogen. The amino acid sequence,
positions of carbohydrate chains and disulfide bridges in the heavy
   SEQUENCE FROM N.A.
MEDLINE-83117859; PubMed-6572010;
Nawa H., Kitamura N., Hirose T., Asai M., Inayama S., Nakanishi S.;
"Primary structures of bovine liver low molecular weight kininogen
   21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
21-JUL-1986 (Rel. 41, Last annotation update)
Kininogen, LAW I precursor (Thiol proteinase inhibitor) (Contains: Bradykinin).
  Bos faurus (Bovine).
Eukaryota: Metazoa; Chordata; Craniata; Vertebrata; Euteleostoml;
Mammalia: Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
Bovidae; Bovinne; Bos.
NCBI_TaxID=9913;
   PIR; A01283; KGBCL1.
InterPro; IRRO00010; Cystatin.
Pfam; PF00031; cystatin; 3.
SMART; SM00043; CY: 3.
PROSITE; PS00287; CYSTATIN; 2.
Glycoprotein; Plasms: Repeat; Vasodilator; Alternative splicing; Thiol protease inhibitor; Bradykinin; Signal.
   CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
PYRROLIDONE CARBOXYLIC ACID.
   N-LINKED (GLCNAC. ..).
O-LINKED (PARTIAL).
   SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
   KININOGEN, LMW I.
   Proc. Natl. Acad. Sci. U.S.A. 80:90-94(1983).
                                  436 AA
   HEAVY CHAIN.
   LIGHT CHAIN.
   BRADYKININ.
  precursors and their two mRNAs.";
   INVOLVED IN BLOOD CLOTTING.
   EMBL; V00426; CAA23709.1; -.
                                  STANDARD;
   378
388
436
  135
257
378
19
87
87
   chain portion.";
J. Biol. Chem. 26
  19
380
389
19
136
19
19
136
                                KNL1_BOVIN
P01046;
  DOMAIN
DOMAIN
MOD_RES
CARBOHYD
CARBOHYD
  CHAIN
PEPTIDE
   CHAIN
  SIGNAL
                       KNL1_BOVIN
  CHAIN
         RESULT
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ö
  Kitamura N., Takagaki Y., Furuto S., Tanaka T., Nawa H., Nakanishi S.; *A single gene for bovine high molecular weight and low molecular weight thingraps **.
  MEDINE-70180420; PubMed-4986212;
Kato H., Nagasawa S., Suzuki T.;
"Studies on the structure of bovine kininogen: cleavages of disulfide
bonds and of methionyl bonds in kininogen-II.";
J. Blochem. 67:313-323(1970).
  Gaps
   Miyata T., Iwanaga S.;
"Bovine high molecular weight kininogen. The amino acid sequence,
positions of carbohydrate chains and disulfide bridges in the heavy
chain portion.";
   MEDLINE-75170265; Pubmed-1169237;
Han Y. N., Komiya M., Iwanaga S., Suzuki T.;
"Studies on the primary structure of bovine high-molecular-weight
"Studies on the primary structure of a fragment ('histidine-rich
peptide') released by plasma kallikrein.";
J. Biochem. 77:55-68(1975).
-!- FUNCTION: (1) KININGENS ARE INHIBITORS OF THIOL PROTEASES; (2)
HMM-KININGEN PLAYS AN IMPORTANT ROLE IN BLOOD COAGULATION BY
HELPING TO POSITION OPTIMALLY PREKALLIKREIN AND FACTOR XI NEXT
   Bradykinin].
Bos taurus (Bovine).
Bukaryotas (Botazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoidea;
Bovidae; Bovinee; Bos
   21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
Kininogen, HMW I precursor (Thiol proteinase inhibitor) [Contains:
   TO FACTOR XII; (3) HMW-KININOGEN INHIBITS THE THROMBIN- AND
   ö
....); OR 169.
  SEQUENCE OF 19-378.
MEDLINE-87137530; PubMed-3546295;
Sueyoshi T., Miyata T., Hashimoto N., Kato H., Hayashida H.,
Miyata T., Iwanaga S.;
   295 A -> T (IN REF. 1; CAA23709).
48427 MW; FO1F7EB6814BCE6C CRC64;
  Length 436;
   3; Indels
N-LINKED (GLCNAC...
N-LINKED (GLCNAC...
N-LINKED (GLCNAC...
INTERCHAIN.
  Score 54; DB 1;
Pred. No. 0.099;
  621 AA
   2; Mismatches
   J. Biol. Chem. 262:2768-2779(1987).
  SEQUENCE FROM N.A. MEDLINE=84014106; PubMed=6571699;
  64.38;
   Nature 305:545-549(1983).
   Conservative
   :: |||||| ||||
292 HDGAFYFKIDTVKKA 306
  STANDARD;
   1 NNATFYFKIDNVKKA 15
168
197
204
406
93
125
1125
124
247
247
266
339
  SECUENCE OF 378-393.
  SEQUENCE OF 458-498
  141
205
228
263
327
350
295
436 AA;
  Query Match
Best Local Similarity
Matches 10; Conserv
   weight kininogens.
   NCBI_TaxID=9913;
```

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EMBL;
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PLASMIN-INDUCED AGGREGATION OF THROMBOCYTES; (4) THE ACTIVE PEPTIDE BRADYKININ THAT IS RELEASED FROM HAW-KININGEN SHOWS A VARIETY OF PHYSIOLOGICAL EFFECTS: (4A) INFUDENCE IN SMOOTH MUSCLE CONTRACTION, (4B) INDUCTION OF HYPOTENSION, (4C) NATRIDERSIS AND DIURESIS, (4D) BECREASE IN BLOOD GLUCOSE LEVEL, (4E) IT IS A MEDIATOR OF INFLAMMATION AND CAUSES (4E1) INCREASE IN VASCULAR PERMERBILITY, (4E2) STIMULATION OF NOCICEPTORS (4E3) RELEASE OF OTHER MEDIATORS OF INFLAMMATION (E.G. PROSTAGLANDINS), (4F) IT HAS A CARRIOPORTECTIVE EFFECT (DIRECTLY VIA BRADYKININ PACTION, INDIRECTLY VIA ENDOTHELIUM-DERIVED RELAXING FACTOR
   SUBCELLULAR LOCATION: Extracellular.
ALTERNATIVE PRODUCTS: 2 isoforms; HMW I (shown here) and LMW I (AC PO1046); are produced by alternative splicing.
TISSUE SPECIFICITY: PLASMA.
PTM: BRADYKININ IS RELEASED FROM KININOGEN BY PLASMA KALLIKREIN.
   CYSTATIN-LIKE 1.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
CYSTATIN-LIKE 3.
PYRROLIDONE CARBOXYLIC ACID.
N-LINKED (GLCNAC. .).
N-LINKED (GLCNAC. .), OR 169.
N-LINKED (GLCNAC. .), OR 169.
N-LINKED (GLCNAC. .), OR 169.
N-LINKED (GLCNAC. .),
  Glycoprotein; Plasma; Repeat; Vasodilator; Alternative splicing; Thiol protease inhibitor; Bradykinin; Blood coagulation; Inflammatory response; Signal.
   Length 621;
  68890 MW; D16850BEFE3C55CD CRC64;
  SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
   KININOGEN, HMW I. HEAVY CHAIN.
   Score 54; DB 1;
Pred. No. 0.14;
  BRADYKININ
  INTERCHAIN
  PIR; A29559; A29559.
InterPro; IPR000010; Cystatin.
InterPro; IPR002395; Kininogen.
Pfam; PF00031; cystatin; 3.
PRINTS; PR00334; KININOGEN.
   PROSITE; PS00287; CYSTATIN; 2.
   64.3%;
66.7%;
  EMBL; V01491; CAA24735.1; -.
   PIR; A01281; KGBOH1.
  621 AA;
  Similarity
  168
197
204
27
82
106
  SM00043;
   DOMAIN
MOD_RES
CARBOHYD
CARBOHYD
  CHAIN
PEPTIDE
   Query Match
   DISULFID
   DISULFID
  SEQUENCE
  CARBOHYD
  CARBOHYD
   CARBOHYD
  DISULFID
  DISULFID
  DISULFID
   DISULFID
  DISULFID
   DISULFID
  DISULFID
   Local
   DOMAIN
  SIGNAL
   DOMAIN
  SMART;
  CHAIN
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LINGUEGRAL CITCHING OI CUNAS IOT MOUSE LOW- and high- molecular kininogen.";

Submitted (APR-1996) to the EMEL/GenBank/DDBJ databases.

-I-FUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)

HEMP-KININOGEN PLAYS AN IMPORTANT ROLE IN BLOOD COAGULATION BY ELLING TO POSITION OPPINALLY PRERALLIKREIN AND FACTOR XI NEXT TO FEACTOR XII: (3) HWW-KININOGEN INHIBITS THE THROMBIN-AND PLASMIN-INDUCED AGGREGATION OF THROMBOLYTES; (4) THE ACTIVE PEPTIDE BRADYKININ THAT IS RELEASED FROM HWW-KININOGEN SHOWS A VARIETY OF PHYSIOLOGICAL EFFECTS: (4A) INFLUENCE IN SMOOTH MUSCLE CONTRACTION, (4B) INDUCTION OF HYPOTENSION, (4C) NATRITURESIS AND DIURESIS, (4D) DECREASE IN BLOOD GLUCOSE LEVEL, (4E) IT IS A MEDIATOR OF INFLAMMATION AND CAUSES (4E1) INCREASE IN VASCULAR PERMEABILITY, (4E2) STIMULATION OF NOCICEPTORS (4E3) RELEASE OF COTHER MEDIATORS OF INFLAMMATION (E.G. PROSTAGLANDINS), (4F) IT HAS A CARDIOPROTECTIVE PEFFORT (DIRECTLY VIA BRADYKININ ACTION), INDIRECTLY VIA BRADYKININ ACTION), CHANNELLY OF STIMULARING (BY SIMILARITY).

CHOWTHAND (BY SIMILARITY).

CHOTTANGEN INHIBITS THE AGGREGATION OF THROMBOCYTES; (6) LAW-KININOGEN NOT INVOLVED IN BLOOD
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  SUBCELLULAR LOCATION: Secreted.
ALTERNATIVE PRODUCTS: 2 ISOFORNS; HWW (SHOWN HERE) AND LAW; ARE PRODUCED BY ALTERNATIVE SPLICING.
TISSUE SPECIFICITY: PLASMA.
PTM: BRADYKININ IS RELEASED FROM KININOGEN BY PLASMA KALLIKREIN.
SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
  PROSITE; PS00287; CYSTATIN; 1. Glycoprotein; Plasma: Repeat; Thiol protease inhibitor; Vasodilator; Bladykinin; Blood coagulation; Inflammatory response; Signal; Alternative splicing.
   Craniata; Vertebrata; Euteleostomi;
Sciurognathi; Muridae; Murinae; Mus.
   STRAIN-C57BL/6 X CBA; TISSUB-Liver;
Takano M., Kondoh J., Yayama K., Okamoto H.;
"Molecular cloning of cDNAs for mouse low- and high- molecular
  HIS-RICH.
INTERCHAIN (BY SIMILARITY).
  KININOGEN.
KININOGEN HEAVY CHAIN.
  KININOGEN LIGHT CHAIN
   16-OCT-2001 (Rel. 40, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
Kininogen precursor [Contains: Bradykinin].
  CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
             661 AA.
   SEQUENCE FROM N.A. (ISOFORMS HWW AND LAWW).
  BRADYKININ.
   POTENTIAL
             PRT;
KNG_MOUSE STANDARD; F
008677; 008676;
16-007-2001 (Rel. 40, Created)
16-007-2001 (Rel. 40, Last sequili-
  MGD; MGT:1097705; Kng.
InterPro; IPR000010; Cystatin.
InterPro; IPR002395; Kininogen.
   Eukaryota; Metazoa; Chordata;
Mammalia; Eutheria; Rodentia;
  EMBL; D84435; BAA19743.1; -.
  D84415; BAA19742.1; -.
   Pfam; PF00031; cystatin; 3.
PRINTS; PR00334; KININOGEN.
SMART; SM00043; CY; 3.
   661
379
388
661
135
257
257
379
524
   Mus musculus (Mouse)
  NCBI_TaxID=10090;
  DOMAIN
DISULFID
  SIGNAL
  DOMAIN
   DOMAIN
  DOMAIN
                THE STATE OF THE S
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Gaps

; 0

3; Indels

Mismatches

5,

Conservative

10;

Matches

:: |||||| |||| HDGAFYFKIDTVKKA 306 1 NNATEYFKIDNVKKA 15

292

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RESULT 9

14

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| ||:||:|
NOKTEFFKLDEFRR 168
                         NNATFYFKIDNVKK
  NUC2_NEUCR
  155
                         -
  Matches
  RESULT 11
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  g
   ð
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  ö
   ö
  N-LINKED (GLCNAC...) (POTENTIAL).
N-LINKED (GLCNAC...) (POTENTIAL).
N-LINKED (GLCNAC...) (POTENTIAL).
VSPPYIAREDERDACTEGEPTHGHOWLHERQ.-> RLLRA
CEYKGRLSKAGAEPAPERQAESSQVKQ (IN ISOFORM
  Gaps
   Gaps
   Baumann L., Baumann P., Moran M.A., Sandstrom J., Thao M.L.;
Genetic characterization of plasmids containing genes encoding
enzymes of leucine biosynthesis in endosymbionts (Buchnera) of tw
related aphid species.";
Submitted (JAN-1998) to the EMBL/GenBank/DBJ databases.
-:- CATALYTIC ACTIVITY: 3-ISOPROPYLMALATE = 2-ISOPROPYLMALEATE +
H(2)0 (ALSO CATALYSES 2-ISOPROPYLMALEATE + H(2)0 = 3-HYDROXY-
  PATHWAY: Leucine biosynthesis; second step.
SUBUNIT: CONSISTS OF TWO DIFFERENT SUBUNITS: LEUC AND LEUD.
   ö
  ;
0
  Score 53; DB 1; Length 661;
Pred. No. 0.22;
2; Mismatches 3; Indels
   48.8%; Score 41; DB 1; Length 208; 42.9%; Pred. No. 7.1; 1:ve 4; Mismatches 4; Indels
   085073;
30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
15-UUN-2002 (Rel. 4). Last annotation update)
3-isopropylmalate dehydratase small subunit (EC 4.2.1.33)
   (Isopropylmalate isomerase) (Alpha-IPM isomerase) (IPMI)
   Leucine biosynthesis; Lyase; Plasmid.
SEQUENCE 208 AA; 24266 MW; E40CD8A63F0396B5 CRC64;
  MISSING (IN ISOFORM LAW).
774460258D58796E CRC64;
   Plasmid pLeu-Dn (pBDn1).
Bacteria; Proteobacteria; gamma subdivision; Buchnera.
   N-LINKED (GLCNAC. . .)
N-LINKED (GLCNAC. . .)
   Buchnera aphidicola (subsp. Diuraphis noxia).
   SIMILARITY.
SIMILARITY.
SIMILARITY.
  SIMILARITY
  SIMILARITY
   SIMILARITY
   4-METHYL-3-CARBOXYPENTANONE).
  EMBL; AF041837; AAD12603.1; -.
InterPro; IPR000573; Aconitase_C.
InterPro; IPR004431; LeuD.
    BY
BY
BY
BY
BY
BY
BY
BY
  Pfam; PF00694; Aconitase_C; 1.
TIGREAMS; TIGR00171; leuD; 1.
  73102 MW;
  63.1%;
66.7%;
  Conservative
   10; Conservative
  |: ||:||| ||||
292 NDHPFYYKIDTVKKA 306
  STANDARD;
  1 NNATFYFKIDNVKKA 15
  433
661 AA;
   Query Match
Best Local Similarity
Matches 6; Conserv
  Best Local Similarity
   NCBI_TaxID=118101;
  SEQUENCE FROM N.A.
  LEUD_BUCDN
  SEQUENCE
  DISULFID
DISULFID
DISULFID
  VARSPLIC
SEQUENCE
DISULFID
  DISULFID
   CARBOHYD
CARBOHYD
   DISULFID
   CARBOHYD
  CARBOHYD
   Query Match
  VARSPLIC
   Matches
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   ö
   Gaps
   STRAIN=74-0R23-IVA;
PUG-24. A Aramayo R., Kang S., Hall J., Metzenberg R.;
NUG-2, a component of the phosphate regulated signal transduction pathway in Neurospora crassa, is an ankyrin repeat protein.";
Dabnitted (MAR-1996) to the EMBL/GenBank/PDBJ databases.
-!- FUNCTION: CONTROLS PHOSPHORUS ACQUISITION.
-!- FUNCTION: CONTROLS 7 ANK REPEATS.
  ö
  Eukaryota; Fungl; Ascomycota; Pezizomycotina; Sordariomycetes;
  Score 41; DB 1; Length 1066;
Pred. No. 38;
   298 ANK 1.
366 ANK 2.
399 ANK 3.
432 ANK 4.
465 ANK 5.
499 ANK 7.
5 117142 MW; 7F672727AFFE92F5 CRC64;
   (Rel. 34, Created)
(Rel. 34, Last sequence update)
(Rel. 34, Last annotation update)
   16-OCT-2001 (Rel. 40, Created)
16-OCT-2001 (Rel. 40, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
Ankyrin repeat protein nuc-2.
  237 AA.
  Pred. No. 38; ; Mismatches
   Sordariales; Sordariaceae; Neurospora
   PROSITE; PS50088; ANK_REPEAT; 4.
PROSITE; PS50297; ANK_REP_REGION; 1.
   EMBL; U51118; AAB03277.1; -.
   48.88;
  70.08;
   InterPro; IPR002110; ANK.
InterPro; IPR004129; GDPD.
   InterPro; IPR004331; SPX.
Pfam; PF03009; ank; 6.
Pfam; PF03009; GDPD; 1.
Pfam; PF03105; SPX; 1.
SMART; SM00248; ANK; 4.
  Conservative
  STANDARD;
  STANDARD;
   370
403
435
470
470
503
1066 AA;
  ANK repeat; Repeat.
REPEAT 268
   Local Similarity
les 7; Conserv
  1 NNATEYFKID 10
   63 NKATFFEQID 72
  SEQUENCE FROM N.A.
  Neurospora crassa
   NCBI_TaxID=5141;
   01-0CT-1996 (
01-0CT-1996 (
01-0CT-1996 (
   YS76_CAEEL
Q09619;
NUC2_NEUCR
Q01317;
   REPEAT
REPEAT
SEQUENCE
  Query Match
  REPEAT
REPEAT
  REPEAT
   REPEAT
  RESULT 12
YS76_CAEEL
```

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Transmembrane;
TRANSMEM 6;
  HFLK_BUCAI
   HFLK_BUCAI
   RESULT 14
        δ
  <del>g</del>
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   RATAILE-21846401; PubMed=11859360;
RA MEDLINE-21846401; PubMed=11859360;
RA WOOD V., Gwilliam R., Rajandream M.A., Lyne M., Lyne R., Stewart A., Squoros J., Peat N., Hayles J., Basham D., Bowman S., As Brooks K., Brown D., Bowman S., Chillingworth T., Churcher C.M., Rateris D., Hidalgo J., Hodgson G., Rentles S., Goble A., Hamlin N., Harris D., Hidalgo J., Hodgson G., Holloyd S., Hornsby T., Howarth S., Huchel E.J., Hunt S., Jagels K., James K., Jones L., Jones M., Laather S., McDonald S., McLean J., RA James K., Jones L., Jones M., Squares S., McDonald S., McLean J., RA Monory P., Moull S., Parison D., Quail M.A., Rabbinowitsch E., Rherford K., Rutter S., Sauders R., Seeger K., Sharp S., Skelton J., Simmonds M., Squares S., Stevens K., Raylor K., Taylor R.G., Parison R., Walsh S.V., Warren T., Whitehead S., Woodward J., Volckaert G., Aert R., Robben J., Grymonprez B., Woltjens I., Vanstreels E., Rieger M., Schaefer M., Mueller-Auer S., Gabel C., Fuchs M., Fritzc C., Holzer E., Moestl D., Hilbert H., Rabbindrik R., Pohl T.M., Berr P., Zimmermann W., Wedler H., Wambutt R., Purnelle B., Goffeau A., Cadleu E., Dreano S., Gloux S., Lelaure V., Mottier S., Adaing Z., Hunt C., Moore K., Hurst S.M., Lucas M., Rochet M., Gaillardin C., Tallada V.A., Garzon A., Thode G., Rar R., Roger M., Galbert F., Rochet M., Gaillardin C., Tallada V.A., Garzon A., Thode G., Rar R., Rochet M., Gaillardin C., Tallada V.A., Garzon A., Phode G., Rar R., Rochet M., Gaillardin C., Tallada V.A., Garzon A., Phode G., Rarga R.R., Cruzado L., Jimenez J., Sanchez M., del Rey F., Benito J.,
  ö
  Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida; Rhabditoidea; Rhabditidae; Peloderinae; Caenorhabditis.
NCBL_TaxID=6239;
  Gaps
  ö
   Score 39; DB 1; Length 237; Pred. No. 18; 2; Indels
Hypothetical 27.3 kDa protein 2K892.6 in chromosome III.
2K892.6.
  Submitted (MAR-1995) to the EMBL/GenBank/DDBJ databases.
  EMBL; Z48638; CAA88568.1; -.
WormPep; ZK892.6; CE01728.
Hypothetical protein.
SEQUENCE 237 AA; 27303 MW; 3C54CD6CF38EA269 CRC64;
   Schizosaccharomyces pombe (Fission yeast).
Kakaryota; Fungi; Ascomycota; Schizosaccharomycetes; Schizosaccharomycetales; Schizosaccharomycetales; Schizosaccharomyces.
   15-JUL-1998 (Rel. 36, Created)
15-JUL-1998 (Rel. 36, Last sequence update)
15-JUL-2002 (Rel. 41, Last annotation update)
19-JUL-1994 (Rel. 41, Last annotation update)
SPAC4F8.04.
  306 AA.
  46.48;
  Query Match
Best Local Similarity 54.5%
---- 6; Conservative
  STANDARD;
                                 Caenorhabditis elegans.
  1 NNATFYFKIDN 11
   60 NDGTFYLKLDS 70
   SEQUENCE FROM N.A. STRAIN-Bristol N2;
  SEQUENCE FROM N.A.
  NCBI_TaxID=4896
   YDS4_SCHPO
014180;
   STRAIN-972;
   Lloyd C.
  RESULT 13
YDS4_SCHPO
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  ö
   Buchnera aphidicola (subsp. Acyrthosiphon pisum) (Acyrthosiphon pisum symbiotic bacterium).
Bacteria: Proteobacteria; gamma subdivision; Buchnera.
  Gaps
  (BY SIMILARITY).
-!- SIMILARITY: BELONGS TO THE BAND 7 / MEC-2 FAMILY. HFLK SUBFAMILY.
Dominguez A., Revuelta J.L., Moreno S., Armstrong J., Forsburg S.L., Cerrutti L., Lowe T., McCombie W.R., Paulsen I., Potashkin J., Shpakovski G.V., Ussery D., Barrell B.G., Nurse P.; The genome sequence of Schizosaccharomyces pombe."; Mature 415:871-880(2002).
-1- SIMILARITY: BELONGS TO THE IMP4 FAMILY.
  -1- SUBUNIT: HFLC AND HFLK MAY INTERACT TO FORM A MULTIMERIC COMPLEX
  Nature 407:81-86(2000).
-1- FUNCTION: HFLC AND HFLK COULD ENCODE OR REGULATE A PROTEASE (BY
  SEQUENCE FROM N.A.
STRAIN=TOKYO 1998;
BDLINE-20445173; PubMed-10993077;
Shigenobu S., Watanabe H., Hattori M., Sakaki Y., Ishikawa H.;
"Genome sequence of the endocellular bacterial symbiont of aphids Buchnera Sp. APS.";
  ö
   46.4%; Score 39; DB 1; Length 306; 37.5%; Pred. No. 23;
  5; Indels
  Hypothetical protein.
SEQUENCE 306 AA; 35835 MW; 2029FBB2358EC01E CRC64;
   P57631;
16-0CT-2001 (Rel. 40, Created)
16-0CT-2001 (Rel. 40, Last sequence update)
16-0CT-2001 (Rel. 40, Last annotation update)
   406 AA.
  5; Mismatches
  POTENTIAL.
  Complete proteome.
7 87 POTE
  EMBL; AP001119; BAB13258.1; -. InterPro; IPR001107; Band.7. InterPro; IPR001972; Stomatin.
  EMBL; 298530; CAB11051.1; -. InterPro; IRP002799; Imp4. Pfam; PF01945; IMP4; 1. Hypothetical protein.
  Pfam; PF01145; Band_7; 1.
PRINTS; PR00721; STOMATIN.
SMART; SM00244; PHB; 1.
   | :||| : |: |:
168 NGPSFYFTLSNLQTAK 183
  6; Conservative
  1 NNATFYFKIDNVKKAR 16
   STANDARD;
   Query Match
Best Local Similarity
Matches 6; Conserv
   NCBI_TaxID=118099
  SIMILARITY).
  Hflk protein.
HFLK OR BU568
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  uterity tour.

11 the transport of   Gaps
   Gaps
  16-0CT-2001 (Rel. 40; Created)
16-0CT-2001 (Rel. 40, Last sequence update)
16-0CT-2001 (Rel. 40, Last annotation update)
Cytidylate kinase (EC 2.7.4.14) (CK) (Cytidine monophosphate kinase)
  Glass J.I., Lefkowitz E.J., Glass J.S., Heiner C.R., Chen E.Y., Cassell G.H.; "The complete sequence of the mucosal pathogen Ureaplasma
   ö
   ö
   Ureaplasma parvum (Ureaplasma urealyticum biotype 1).
Bacteria; Firmicutes; Mollicutes; Mycoplasmataceae; Ureaplasma
NCBL_TaxID=134821;
  45.2%; Score 38; DB 1; Length 224; 66.7%; Pred. No. 25;
  Score 39; DB 1; Length 406;
Pred. No. 31;
2; Mismatches 3; Indels
  4; Indels
  InterPro; IPR003136; Cytidylate_kin.
Pfam; PF02224; Cytidylate_kin. 1.
TIGRFAMS; TIGR00017; cmk; 1.
Transferase; Kinase; ATP-binding; Complete proteome.
NP_BIND 11 19 ATP (BY SIMILARITY).
SEQUENCE 224 AA; 25618 MW; AEEF7DFF423DADFF CRC64;
406 AA; 46928 MW; 70BA5DD72E537DB2 CRC64;
  224 AA
  0; Mismatches
  PRT;
   STRAIN-Serovar 3;
MEDLINE-20500219; Pubmed-11048724;
  EMBL; AE002131; AAF30751.1; -. HSSP; P23863; 1CKE.
   46.48;
  8; Conservative
   Conservative
  STANDARD;
  2 NATFYFKIDNVK 13
   334 NNPMFFFSLDN 344
   Query Match
Best Local Similarity
   1 NNATEYFKIDN 11
   Best Local Similarity
Matches 8; Conserv
  SEQUENCE FROM N.A.
  (CMP kinase)
CMK OR UU342
   urealyticum.
   9
   KCY_UREPA
Q9PQE9;
SEQUENCE
  Query Match
   RESULT 15
KCY_UREPA
   Matches
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Search completed: March 13, 2003, 19:13:26 Job time: 12 secs THIS PAGE BLANK (USPTO)

| GenCore version 5.1.4_p5_4578<br>Copyright (c) 1993 - 2003 Compugen Ltd.                                            |      |
|---------------------------------------------------------------------------------------------------------------------|------|
| OM protein - protein search, using sw model                                                                         |      |
| Run on: March 13, 2003, 19:07:27 ; Search time 33.3333 Seconds (without alignments) 98.903 Million cell updates/sec | •    |
| Title: US-09-461-061A-10 Perfect score: 84 Sequence: 1 NNATFYFKIDNVKKAR 16                                          |      |
| Scoring table: BLOSUM62<br>Gapop 10.0 , Gapext 0.5                                                                  |      |
| Searched: 671580 seqs, 206047115 residues                                                                           |      |
| Total number of hits satisfying chosen parameters: 671580                                                           |      |
| Minimum DB seq length: 0<br>Maximum DB seq length: 200000000                                                        |      |
| Post-processing: Minimum Match 0%<br>Maximum Match 100%<br>Listing first 45 summaries                               | ·· : |
| Database : SPTREMBL_21:*                                                                                            |      |

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution. sp\_archea:\*
sp\_bacteria:\*
sp\_fungi:\*
sp\_human:\*
sp\_invertebrate:\*
sp\_mammal:\*
sp\_phage:\*
sp\_phage:\*
sp\_plant:\*
sp\_plant:\*
sp\_rodent:\*
sp\_vartebrate:\*


| Description                   | P70517 rattus norv 063581 rattus norv 061xk5 mus musculu Q84233 dictyosteli Q9k101 porphyromon G9807 listeria mo Q8w112 chlorogoniu Q8w111 ankistrodes Q9zut5 arabidopsis 029973 archaeeglub Q55p10 dictyosteli Q8xmh6 clostridium Q8xmh6 clostridium Q8xmh6 clostridium Q8xmh6 pyrococcus Q941z phaseolus v Q9emr4 amsacta moo                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
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| SUMMARIES                     | P70517<br>Q63581<br>Q63581<br>Q612X5<br>Q812A3<br>Q871B1<br>Q87807<br>Q8W112<br>Q98W11<br>Q92UT5<br>Q95P10<br>Q8XM46<br>Q98XM6<br>Q94V021<br>Q94 IR2                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| DB                            | 111<br>111<br>111<br>110<br>110<br>110<br>110<br>110                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| å<br>Query<br>Match Length DB | 423<br>423<br>432<br>806<br>806<br>866<br>529<br>229<br>229<br>154<br>657<br>467<br>467<br>467<br>467<br>469<br>873<br>873                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| &<br>Query<br>Match           | 666.77<br>666.77<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>667.71<br>66 |
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| 098GP2<br>0017098<br>0045699<br>009GRX5<br>099RF1<br>086664<br>087KU8<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7<br>097RD7                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | ALIGNMENTS  PRT; 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| 55<br>55<br>56<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57<br>57                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | 02,<br>02,<br>020,<br>11 pu<br>120,<br>120,<br>13-25,<br>04, 17<br>04, 17<br>0                                           |
| 996<br>1001<br>1546<br>15566<br>15676<br>15676<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>16776<br>1677 | MINARY, BLrel. BLrel. alpha (alpha) (a) Choole (b) Choole (c) Choo                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
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MEDLINE-20499374; PubMed-11042159;
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STRAIN-C57BL/6J; TISSUE-PLACENTA;
Adachi J., Aizawa K., Akahira S., Akimura T., Aono H., Arai A.,
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01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
01-JUN-Emalle placenta cDNA, RIKEN full-length enriched library,
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  01-NOV-1996 (TrEMBLrel. 01, Created)
01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-MAR-2002 (TrEMBLrel. 20, Last annotation update)
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  Created)
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STRAIN-C57BL/64; TISSUE-PLACENTA; MEDLINE=20530913; PubMed=11076861; Shibata K., Toh M., Alzawa K., Naqaoka S., Sasaki N., Carninci P., Shibata K., Toh M., Alzawa K., Naqaoka S., Sasaki N., Carninci P., Sumi N., Ishii Y., Nakamura S., Hazama M., Nishine T., Tashiro H., Itoh M., Yamanoto R., Matsumoto H., Sakaguchi S., Ikegami T., Kashiwagi K., Fujiwake S., Inoue K., Togawa Y., Izawa M., Ohara E., Watshiwagi K., Fujiwake S., Ishikawa T., Ozawa K., Tanaka T., Matsuura S., Kawai J., Okazaki Y., Muramatsu M., Inoue Y., Kira A., Hayashizaki Y.; Firk integrated sequence analysis (RISA) system-384-format Genome Res. 10:1757-1771(2000).
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  Score 53; DB 11; Length 432;
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Submitted (DEC-2001) to the EMBL/GenBank/DDBJ databases.
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Pfam; PF00031; Cystatin; 3.
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   Gloeckner G., Elchinger L., Szafranski K., Pachebat J., Dear P., Lehmann R., Baumgart C., Parra G., April J.F., Guigo R., Kumpf K., Tunggal B., Cox E., Quail M.A., Platzer M., Rosenthal A., Noegel A.A.; "Sequence and Analysis of Chromosome 2 of Dictyostelium."; Submitted (MAR-2002) to the EMBL/GenBank/DDBJ databases.

EMBL; AC115678; AAL92605.1; -
  Ross B.C.;
"Genomic analysis of Porphyromonas gingivalis for vaccine discovery.";
"Genomic analysis of Porphyromonas gingivalis for vaccine discovery.";
Submitted (FEB-2000) to the EMBL/GenBank/DDBJ databases.
EMBL; AF237557; AAF81415.1;
Interpro; IPR000531; TONB_DOXC.
Interpro; IPR000531; TONB_DOXC.
  01-0CT-2000 (TTEMBLrel. 15, Created)
01-0CT-2000 (TTEMBLrel. 15, Last sequence update)
01-DCC-2001 (TTEMBL-11), Last annotation update)
Hypochetical tonB-linked outer membrane receptor PG47.
Porphyromonas gingivalis (Bacteroides gingivalis).
Bacteria; CFB group; Bacteroidetes; Bacteroidales; Porphyromonadaceae;
  Gaps
   Gaps
  Σ.
  ö
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   Ross B., Barr I., Patterson M., Agius C., Rothel L., Margetts Hocking D., Webb E.;
Hocking D., Webb E.;
"P. gingivalis polypeptides and nucleic acids.";
Submitted (FEB-2000) to the EMBL/GenBank/DDBJ databases.
   Score 44; DB 2; Length 867;
Pred. No. 93;
2; Mismatches 5; Indels
  53.6%; Score 45; DB 5; Length 806; 57.1%; Pred. No. 60;
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SEQUENCE 806 AA; 96216 MW; 2D9BDB77A5BAD1F6 CRC64;
  867 AA; 95231 MW; FAB4842E821C4E8E CRC64;
   01-JUN-2002 (TrEMBLrel. 21, Created)
01-JUN-2002 (TrEMBLrel. 21, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
D1-JUN-2002 (TrEMBLrel. 21, Last annotation update)
UCBI_TaxID-44669;
  466 AA.
  867 AA.
  Pred. No. 60;
1; Mismatches
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  52.4%;
53.3%;
   Query Match
Best Local Similarity 53.3.
Local 8; Conservative
  Best Local Similarity 57.1
Matches 8; Conservative
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816 STTFYFNMDNVADKR 830
  PRELIMINARY;
  PRELIMINARY;
  146 NNYPFYFTLDETKK 159
   2 NATFYFKIDNVKKAR 16
   1 NNATFYFKIDNVKK 14
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  SEQUENCE FROM N.A.
  SEQUENCE FROM N.A.
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NCBI_TaxID=837;
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   STRAIN-AX4;
  STRAIN-W50;
  Receptor:
  SEQUENCE
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   Q8Y807
Q8Y807;
   09KIB1;
   Q9KIB1
   RESULT 5
Q9KIB1
  RESULT 6
Q8Y807
ID Q8Y80
AC Q8Y80
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Glaser P., Frangeul L., Buchteser C., Rusniok C., Amend A.,
A Glaser P., Frangeul L., Buchteser C., Rusniok C., Amend A.,
Charbit A., Chetouani F., Couve B., de Daruvar A., Dehoux P.,
Domann E., Dominguez-Bernal G., Duchaud E., Durant L., Dussurget O.,
Entian K.-D., Fsihi H., Garcia-del Portillo F., Garrido P.,
A Gutter L., Goebel W., Gomez-Lopez N., Hain T., Hauf J., Jackson D.,
Jones L.-M., Kaerst U., Kreft J., Kuhn M., Kunst F., Kurapkat G.,
Madueno E., Maltournam A., Mata Vicente J., Ng E., Nedjari H.,
Nordsiek G., Novella S., de Pablos B., Perez-Diaz J.-C., Purcell R.,
Remmel B., Rose M., Schlueter T., Simosa N., Tierrez A.,
Vazquez-Boland J.-A., Voss H., Wehland J., Cossart P.;
Comparative genomics of Listeria species.";
EMBL; AL591977; CAC99190.1;
Listilist; LMO01112;
Listilist; LMO01113;
Listilist; LMO01112;
Listilist; LMO01112;
Listilist; LMO01112;
Listilist; LMO01112;
Listilist; LMO01112;
Listilist; LMO01112;

   SEQUENCE FROM N.A.

MEDLINE=97331047; PubMed=9185572;

Turmel M. Otis C., Cote V., Lemieux C.;

Turmel M. Otis C., Cote V., Lemieux C.;

Turmel M. Otis C., Cote V., Lemieux C.;

I-Ceui homing endonuclease.";

Nucleic Acids Res. 25:2610-2619(1997).

EMBL: L42860, AAL3436.1; -.

Pfam: PF00961; LAGLIDADG_1: 1.

Chloroplast.
   Gaps
  Gaps
   01-MAR-2002 (TrEMBLrel. 20, Created)
01-MAR-2002 (TrEMBLrel. 20, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
Chloroplast large-subunit ribosomal RNA (rrnL), site-specific DNA endonuclease I-Cell gene.
Chlorogonium elongatum.
  Eukaryota Viridiplantae; Chlorophyta; Chlorophyceae; Volvocales; Haematococcaceae; Chlorogonium.
NCBL_TaxID=52029;
   ö
   ö
   Listeria monocytogenes.
Bacteria; Firmicutes; Bacillus/Clostridium group; Bacillales;
Listeriaceae; Listeria.
NCBL_TaxID=1639;
  Score 43; DB 16; Length 466;
Pred. No. 74;
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   Phram, PPO1580; FtsK.SpolIIE; 1.
Hypothetical protein; Complete proteome.
SEQUENCE 466 AA; 53666 MW; ED1937AD833536F3 CRC64;
   229 AA; 26622 MW; 5C7EF3D0DC27521E CRC64;
(TrEMBLrel. 20, Created)
(TrEMBLrel. 20, Last sequence update)
(TrEMBLrel. 21, Last annotation update)
   8
   229 AA
  50.0%; Score 42; DB 72.7%; Pred. No. 54; ive 1; Mismatches
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  SEQUENCE FROM N.A.
STRAIN-EGD-E / SEROVAR 1/2A;
MEDLINE-21537279; PubMed-11679669;
   Hypothetical protein lmol112.
   51.2%;
60.0%;
  Ouery Match
Best Local Similarity 60.v.
   Conservative
  PRELIMINARY;
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RESULT 8 Q8WL11

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Klenk H.-P., Clayfon R.A., Tomb J.-F., White O., Nelson K.E., Ketchum K.A., Dodson R.J., Gwinn M., Hickey E.K., Peterson J.D., Richardson D.L., Kerlavage A.R., Graham D.E., Kyrpides N.C., Eleischmann R.D., Quackenburb J., Lee N.H., Sutton G.G., Gill S., Kirkness E.F., Dougherty B.A., McKenney K., Adams M.D., Loftus B., Verbeek R., Gocayne J.D., Weidman J.F., McDonald L., Utterback T., Cotton M.D., Spriggs T., Artiach P., Kaine B.P., Sykes S.M., Sadow P.W., D'Andrea R.P., Bowman C., Fujil C., Garland S.A., Wenter J.C., Venter J.C., Kinse B.P., Sykes S.M., State J.C., Saith H.O., Woese C.R.,
   "The complete genome sequence of the hyperthermophilic, sulphate-reducing archaeon Archaeoglobus fulgidus.";
Nature 390:364-370(1997).
  Score 41.5; DB 17; Length 469;
Pred. No. 1.3e+02;
  Length 657;
  3; Indels
  1; Indels
Submitted (MAR-2000) to the EMBL/GenBank/DDBJ databases.
EMBL; AC005896; AAC98051.1; -.
InterPro; IPR002885; PPR.
Pfam; PF01555; PPR, 13.
TIGRFAMS; TIGR00756; PPR; 12.
SEQUENCE 657 AA; 73342 MW; 9F76D544896EB872 CRC64;
   Archaea; Buryarchaeota; Archaeoglobi; Archaeoglobales; Archaeoglobaceae; Archaeoglobus.
NCBI_TaxID=2234;
  469 AA; 51663 MW; 3BDAA6AFB179BB43 CRC64;
   DHKF.
Dictyostelium discoideum (Slime mold).
Eukaryota: Mycetozoa; Dictyosteliida; Dictyostelium.
NCBL_max1D+44669;
   01-JAN-1998 (TrEMBLrel. 05, Created)
01-JAN-1998 (TrEMBLrel. 05, Last sequence update)
01-MAR-2002 (TrEMBLrel. 20, Last annotation update)
Hypothetical protein AF1231.
  Last sequence update)
Last annotation update)
  50.0%; Score 42; DB 10;
53.8%; Pred. No. 1.5e+02;
iive 3; Mismatches 3
  469 AA
  PRT; 1548 AA
  83.3%; Pred. no.
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  SEQUENCE FROM N.A.
STRAIN-VC-16 / DSM 4304 / ATCC 49558;
MEDLINE-98049343; PubMed-9389475;
   01-DEC-2001 (TrEMBLrel. 19, Created)
01-UNW-2002 (TrEMBLrel. 21, Last seq
01-UNW-2002 (TrEMBLrel. 21, Last and
Histidine kinase DMKF.
   EMBL; AE001019; AAB90020.1; -. IIGR; AF1231; -.
  49.48;
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Matches 7; Conservative
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   4 TFYFKIDNVKKAR 16
  335 NATFYLKI-NVK 345
   2 NATFYFKIDNVK 13
   Hypothetical
  Query Match
  SEQUENCE
  Query Match
  Q95PIO;
  029037
   095PI0
  RESULT 10
029037
  RESULT 11
Q95PI0
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   SEQUENCE FROM N.A.

STRAIN-CV. COLUMBIA;

WEDLINE-20083487; PubMed=10617197;

Lin X., Kaul S., Rounsley S.D., Shea T.P., Benito M.-I., Town C.D.,

Fujii C.Y., Mason T.M., Bowman C.L., Barnstead M.E., Feldblyum T.V.,

Buell C.R., Ketchum K.A., Lee J.J., Ronning C.M., Koo H., Moffat K.S.,

Cronin L.A., Shen M., VanAken S.E., Umayam L., Tallon L.J., Gill J.E.,

Adams M.D., Carrera A.J., Creasy T.H., Goodman H.M., Somerville C.R.,

Copenhaver G.P., Preuss D., Nierman W.C., White O., Eisen J.A.,

Salzberg S.L., Fraser C.M., Venter J.C.;

Sequence and analysis of chromosome 2 of the plant Arabidopsis
   Arabidopsis thaliana (Mouse-ear cress).

Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
eurosids II; Brassicales; Brassicaceae; Arabidopsis.
   Eukaryota; Viridiplantae; Chlorophyta; Chlorophyceae; Selenastraceae;
  MEDIINE-97331047; Pubmed-9185572;
Turmel M., Otis C., Cote V., Lemieux C.;
"Evolutionarily conserved and functionally important residues in the
  Gaps
   01-MAR-2002 (TrEMBLrel. 20, Created)
01-MAR-2002 (TrEMBLrel. 20, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
Chloroplast large-subunit ribosomal RNA (rrnL), site-specific DNA
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  Score 42; DB 8; Length 244;
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  Last sequence update)
Last annotation update)
  244 AA.
   657 AA.
   1; Mismatches
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Nucleic Acids Res. 25:2610-2619(1997).
EMBL; L42984; AAL34330.1; -.
InterPro; IPR001982; Endonuc_LAG/HNH.
Pfam; PF00961; LAGLIDADG_1; 1.
   Created)
  PRT;
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Best Local Similarity /2...
**Local Similarity /2...
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  PRELIMINARY;
   PRELIMINARY;
  Nature 402:761-768(1999)
                       :||| |||||
126 SNATMVFKIDN 136
   130 SNATFVFTIDN 140
1 NNATEYFKIDN 11
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Lin X.;
   1 NNATEYFKIDN 11
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   SEQUENCE FROM N.A
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   Ankistrodesmus.
   Chloroplast.
   Chloroplast.
  SEQUENCE
   Q9ZUT5
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RESULT 9 Q9ZUT5

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Gaps

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P SEQUENCE FROM N.A.

P SEQUENCE FROM N.A.

STRAIN-ATCZ 25586;

X MEDLINE-21886394; PubMed-11889109;

K Rapatral V., Anderson I., Ivanova N., Reznik G., Los T., Lykidis A.,

A Bhattacharyya A., Bartman A., Gardner W., Grechkin G., Zhu L.,

A Bhattacharyya A., Bartman A., Gardner W., Grechkin G., Bernal A.,

Larsen N., D'Souza M., Walunas T., Pusch G., Haselkorn R.,

T "Genome sequence and analysis of the oral bacterium Fusobacterium nucleatum strain ATCZ 25586.";

J. Bacteriol. 184:2005-2018(2002).
  "Pyrococcus abyssi genome sequence: insights into archaeal chromosome
  Archaea; Euryarchaeota; Thermococci; Thermococcales; Thermococcaceae;
   Gaps
   Gaps
  Phaseolus vulgaris (Kidney bean) (French bean).
Eukaryota, Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; eudicotyledons; core eudicots; Rosidae;
eurosids I; Fabales; Fabaceae; Papilionoideae; Phaseoleae; Phaseolus
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0
   ö
   Length 454;
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   Score 41; DB 16; Length 41.
Pred. No. 1.4e+02;
   Indels
   structure and evolution.";
Submitted (JUL-1999) to the EMBL/GenBank/DDBJ databases
   EMBL; AJ24885; CAB49560.1; -.
InterPro; IPR000357; HEAT_repeat.
PROSITE; PS50077; HEAT_REPEAT; 1.
Hypothetical protein; Complete processes.
SEQUENCE 454 Aa; 52171 MW; B44FB5C2C7919936 CRC64;
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  01-MAY-2000 (TrEMBLrel. 13, Created)
01-MAY-2000 (TrEMBLrel. 13, Last sequence update)
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Carotenoid 9,10-9',10' cleavage dioxygenase.
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Pred. No. 1.5e+02;
   454 AA
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PAB1940.
   48.8%;
57.1%;
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316 SNATSYIKIDDAMK 329
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  Pyrococcus abyssi
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  Heilig R.
  09V0Z1;
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  094IR2
  Q9V0Z1
   datches
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   RESULT 15
   RESULT 14
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  Q9V0Z1
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   Gaps
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   Shimizu T., Ohtani K., Hirakawa H., Ohshima K., Yamashita A., Shiba T., Ogasawara N., Hattori M., Kuhara S., Hayashi H.; "Complete genome sequence of Clostridium perfringens, an anaerobic
   Anjard C., Loomis W.F.;
"The histidine kinases of Dictyostelium.";
(In) Inouye M., Dutta R. (eds.);
Histidine Kinases in Signal Transduction, pp.1-1, Academic Press,
   ;
   ó
  Score 41.5; DB 5; Length 1548;
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   48.8%; Score 41; DB 16; Length 154;
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   Anjard C., Loomis W.F.;
Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases
EMBL; AF362368; AAK54087.2; -.
   Proc. Natl. Acad. Sci. U.S.A. 99:996-1001(2002).
EMBL. AP003187; BAB80419.11, -
Hypothetical protein; Complete proteome.
SEQUENCE 154 AA; 17971 MW; 97C25B68403A4B5B CRC64;
   OSRIGS;
01-JUN-2002 (TrEMBLrel. 21, Created)
01-JUN-2002 (TrEMBLrel. 21, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
   01-MAR-2002 (TTEMBLrel. 20, Created)
01-MAR-2002 (TTEMBLrel. 20, Last sequence update)
01-MAR-2002 (TTEMBLrel. 20, Last annotation update)
Hypothetical protein CPE0713.
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Bacteria; Fusobacteria; Fusobacterium.
NCBI_TaxID=76856;
   3; Mismatches
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  PRT;
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   49.4%;
   53.8%;
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Best Local Similarity 60.0.
   Best Local Similarity 53.8
Matches 7; Conservative
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  PRELIMINARY;
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Clostridium perfringens.
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   | |||:|:||:||:
| 128 TLYFKLDYLKKKK 140
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   San Diego (2001).
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   Kinase.
  9нжжө
  08кн68
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Q8RH68
  RESULT 12
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NCBI\_TaxID=3885;

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   0; Gaps
                           Schwartz S.H., Qin X., Zeevaart J.A.D.;
Schwartz S.H., Qin X., Zeevaart J.A.D.;
Schwartz S.H., Qin X., Zeevaart J.A.D.;
Schwartz S.H., Qin X., Zeevaart J.A.D.;
Submitted (APR-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; ARC02525; ARX3844.1;
Interpro; IPR004294; RPE65.
Pfam; PP03055; RPE65.
Dioxygenase.
SQUENCE 543 AA; 61100 MW; F22C9883A05325F7 CRC64;
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298 NKTLIFSFDSTKKAR 312
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GenCore version 5.1.4_p5_4578 Copyright (c) 1993 - 2003 Compugen Ltd.
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| OM protein - protein search, using sw model | March 13, 2003, 19:07:07; search timé 21 seconds (without alignments) 50.762 Million cell updates/sec |
| OM protei                                   | Run on:                                                                                               |

| A-1             |                |              |  |
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| US-09-461-061A- | 45             | 1 NNATFYFK 8 |  |
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| Scoring  |            |

<sup>908470</sup> seqs, 133250620 residues Searched:

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|--------------------------------|------------|--------------|
| Match                          | Match 100% | first 4      |
| Minimum                        | Maximum    | Listing      |
| Post-processing: Minimum Match |            |              |

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|-------|-------------------------------------------------------------------|
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| 7:    | /SIDS1/gcgdata/geneseq/geneseqp-emb1/AA1986.DA1                   |
| <br>æ | /SIDS1/gcgdata/geneseq/geneseqp-embl/AA1987.DA1                   |
| 6     |                                                                   |
| 10:   |                                                                   |
| 11:   |                                                                   |
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| 13:   |                                                                   |
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<sup>/</sup>SIDS1/gcgdata/geneseq/geneseqp-embl/AA1998.DAT /SIDS1/gcgdata/geneseq/geneseqp-embl/AA1999.DAT /SIDS1/gcgdata/geneseq/geneseqp-embl/AA2000.DAT /SIDS1/gcgdata/geneseq/geneseqp-embl/AA2000.DAT /SIDS1/gcgdata/geneseq/geneseqp-embl/AA2001.DAT /SIDS1/gcgdata/geneseq/geneseqp-embl/AA2002.DAT 114. 115:: 117:: 120:: 221:: 23::

/gcgdata/geneseq/geneseqp-emb1/AA1996.DAT

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## Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

| 3                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |            |
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| 1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | CHICANOTTO |
| ;                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | o the      |
| )                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |            |
| profit properties of the second secon |            |
| 7                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |            |
| }                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |            |

| Description                       | Human kininogen D3 | Anti-angiogenic D3 | Anti-angiogenic D3 | Anti-angiogenic D3 | Anti-anglogenic D3 | Domaine 3, bradyki | Human kininogen D3 | Human high mol.wt. | Novel human diagno | Novel human diagno |
|-----------------------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|
| ΙD                                | AAB37455           | AAY95405           | AAY95409           | AAY95410           | AAY95408           | AAR33350           | AAB37447           | AAY95426           | ABG21102           | ABG21.099          |
| DB                                | 21                 | 21                 | 21                 | 21                 | 21                 | 14                 | 21                 | 21                 | 22                 | 22                 |
| Length                            | 6                  | 10                 | 16                 | 16                 | 32                 | 117                | 122                | 123                | 248                | 369                |
| %<br>Query<br>e Match Length DB I | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              | 100.0              |
| Score                             | 45                 | 45                 | 45                 | 45                 | 45                 | 45                 | 4.5                | 45                 | 45                 | 45                 |
| Result<br>No.                     | г                  | 7                  | m                  | *                  | 2                  | 9                  | 7                  | 80                 | თ                  | 10                 |

| Human high molecul |              | high    | Bradykinin analogo | poly   |        | Novel human connec | Lactococcus lactis | Staphylococcus hom | ia     | Bacillus sp. GUS p | sp. Gus | sp. codo | Bacillus sp. GUS p | Staphylococcus bet | Codon-optimised St |            | ne/haema | ιn     | ED     | Ø    | Drosophila melanog | Human secreted pro | Human immune/haema | Human secreted pro | Human mutant cysta | Human mutant cysta | Protein which is s | Human ORFX protein |            | Novel human diagno | Macaque platelet a | Macaque platelet-a |        | macaque      |  |
|--------------------|--------------|---------|--------------------|--------|--------|--------------------|--------------------|--------------------|--------|--------------------|---------|----------|--------------------|--------------------|--------------------|------------|----------|--------|--------|------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|------------|--------------------|--------------------|--------------------|--------|--------------|--|
|                    | 22           | 23      | 18 AAW5433         | 22     | 22     | 22                 |                    | 21                 | 22     | 20                 | 50      | 20       | 50                 | 21                 | 21                 | 20 AAW9382 | 22       | 21     |        | 21   | 22                 |                    | 22                 | 21                 | 21                 | 21 AAY8118         | 50                 | 23                 | 22 AA01296 | 22                 | _                  | 18                 | 18     | <del>-</del> |  |
| 100.0              | 45 100.0 644 | 5 100.0 | 9 86.7             | 0.08 9 | 4 75.6 | 4 75.6             | 4 75.6             | 4 75.6             | 4 75.6 | 75.6               | 4 75.6  | 4 75.6   | 4 75.6             | 4 75.6             | 4 75.6             | 4 75.6     | 3 73,3   | 3 73.3 | 3 73.3 | 73.3 | 2 71.1 1           | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9     | 1 68.9             | 1 68.9             | 1 68.9             | 1 68.9 | 1 68.9       |  |
| 11                 | 12           | 13      | 14                 | 15     | 16     | 17                 | 18                 | 19                 | 20     | 21                 | 22      | 23       | 24                 | 25                 | 26                 | . 27       | 28       | 29     | 30     | 31   | 32                 | 33                 | 34                 | 35                 | 36                 | 37                 | 3.8                | 39                 | 40         | 41                 | 42                 | 43                 | 44     | 45           |  |

## ALIGNMENTS

|        |          |                                   |   |           |   |                           |    | .:                                  |   |
|--------|----------|-----------------------------------|---|-----------|---|---------------------------|----|-------------------------------------|---|
|        |          | AA.                               |   |           |   |                           |    | agment                              |   |
|        |          | σ                                 |   |           |   |                           |    | fr                                  |   |
|        |          | AAB37455 standard; peptide; 9 AA. |   |           |   | 21-FEB-2001 (first entry) |    | Human kininogen D3 peptide fragment |   |
|        |          | ġ                                 |   |           |   | rst                       |    | D3                                  |   |
|        |          | dar                               |   |           |   | (£1                       |    | en                                  |   |
|        |          | tan                               |   |           |   | ᅼ                         |    | nog.                                |   |
|        |          | 55                                |   | .,        |   | 200                       |    | ini                                 |   |
|        |          | 745                               |   | 745       |   | ÈB-                       |    | =                                   |   |
| -      | 55       | AB3                               |   | AAB37455; |   | 1-E                       |    | luma                                |   |
| Ξ      | AAB37455 | ⋖                                 |   | Æ         |   | N                         |    | Æ                                   |   |
| RESULT | AAB      | G                                 | X | AC        | × | 덥                         | XX | DE                                  | × |
|        |          |                                   |   |           |   |                           |    |                                     |   |

Enzyme; legumain; endopeptidase; cystatin; human; kininogen. XXX

|               |                 |              | 00WO-GB01571.               |
|---------------|-----------------|--------------|-----------------------------|
| Homo sapiens. | WO200064945-A1. | 02-NOV-2000. | 20-APR-2000; 2000WO-GB01571 |
| SO            | X Z 3           | ₹ <b>2</b> ; | PF                          |

99GB-0009133. Abrahamson M, Barrett AJ; (BABR-) BABRAHAM INST. 22-APR-1999; 

WPI; 2000-687316/67.

Inhibition of mammalian legumain or legumain-related endopeptidase by cystatin involves interaction with second papain-non-reactive site of cystatin .

Disclosure; Fig 4; 45pp; English.

<sup>908470</sup> Total number of hits satisfying chosen parameters:

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Claim 1; Page 25; 44pp; English.
        \overset{\circ}{\mathbf{x}}\overset{\circ}
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О
   legumain or a legumain-related endopeptidase by cystatin. The inhibition involves an interaction between legumain and a papain-non-reactive site of cystatin. Legumain (EC 3.4.22.34) is a cysteine endopeptidase, and performs a protein-processing function. The present sequence is a peptide fragment of human kininogen D3, which was used in the present invention. Kininogen is a type 3 cystatin. The present sequence is thought to be involved in a legumain-inhibitory site.
   ö
  Composition for inhibiting angiogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rheumatoid arthritis, and ocular disorders comprises a kininogen domain
   Gaps
   Anti-anglogenic; anglogenesis; inhibitor; kininogen; homologue; endothelial cell proliferation; apoptosis; cancer; ocular disorder; rheumatoid arthritis; cytostatic; antiarthritic; antirheumatic; therapy; human; D3 peptide.
  "corresponds to residues Asn(275)-Lys(282) of HK domain 3"
   sequence of AAY95407 or its C-terminal truncation fragment containing at least 1 amino acid"
   sequence of AAY95406 or its N-terminal truncation fragment containing at least 1 amino acid"
   ö
   "Xaa represents 0 amino acids, or the
  Length 9;
   Indels
  "Xaa represents 0 amino acids,
                                    present invention relates to inhibition of the
  100.0%; Score 45; DB 21;
Similarity 100.0%; Pred. No. 7.8e+05;
8; Conservative 0; Mismatches 0;
   Location/Qualifiers
  AAY95405 standard; Peptide; 10 AA.
   98US-0112427.
  99WO-US28465
  Anti-angiogenic D3 peptide.
   (first entry)
  /note=
  /note=
   /note=
  WPI; 2000-442247/38.
  (UTEM ) UNIV TEMPLE.
  Best Local Similarity
Matches 8; Conserv
  (MCCR/) MCCRAE R K.
   Misc-difference 10
   9
A
   Key
Misc-difference
  1 NNATEYEK 8
   WO200035407-A2
   25-SEP-2000
   Homo sapiens
   16-DEC-1998;
  02-DEC-1999;
  22-JUN-2000.
   RK;
   AAY95405;
   Sequence
  Query Match
  analog
  Peptide
   McCrae
  AAY95405
   ò
  g
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ö
The present sequence is that of a D3 peptide derived from high mol.wt. kininogen (HK) domain 3 (see AAY95426). The D3 peptide, which may optionally include N terminal and/or C-terminal protecting groups, inhibits endothelial cell proliferation and thus possesses anti-angiogenic activity. It is an example of peptides of the invention (see AAY95405-26) which are analogues of certain sites in the HK domain 3, in this case amino acids ABAD75-Lys282. The peptides inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including such peptides are used in claimed methods for inhibiting anglogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell proliferation, and inducing endothelial cell proliferation, and inducing endothelial cell cancer, rheumatoid arthritis, and ocular disorders characterized by undesired vascularization of the retina are treated.
   Composition for inhibiting angiogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rhewmatoid arthritis, and ocular disorders comprises a kininogen domain
   The present sequence is that of a D3 peptide derived from human high mol.wt. kininogen (HK) domain 3 (see AAY95426). The D3 peptide inhibits endothelial cell proliferation and thus possesses anti-anglogenic activity. It is an example of D3 peptides of the invention (see AAY95405-26) that are analogues of certain sites in the HK domain 3, in this case amino acid residues Asn275-Lys282. The peptides inhibit endothelial cell proliferation and may also
   Gaps
  Anti-angiogenic; angiogenesis; inhibitor; kininogen; homologue; endothelial cell proliferation; apoptosis; cancer; ocular disorder; rheumatoid arthritis; cytostatic; antiarthritic; antirheumatic;
  ;
0
   DB 21; Length 10;
   Indels
   0.061;
  Score 45; DBPred. No. 0.06Mismatches
  ¥
   Claim 6; Page 26; 44pp; English.
   AAY95409 standard; Peptide; 16
  100.0%;
   99WO-US28465
   98US-0112427
  Anti-angiogenic D3 peptide
  (first entry)
   therapy; human; D3 peptide
  Query Match 100.
Best Local Similarity 100.
Matches 8; Conservative
   WPI; 2000-442247/38.
  (UTEM ) UNIV TEMPLE
   (MCCR/) MCCRAE R K.
  10 AA;
  œ
  σ
   WO200035407-A2
   1 NNATFYFK
  Homo sapiens.
   02-DEC-1999;
   16-DEC-1998;
  25-SEP-2000
  22-JUN-2000.
   McCrae RK;
   Sequence
   AAY95409;
  analog
   AAY95409
   RESULT
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Gaps

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Composition for inhibiting angiogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rheumatoid arthritis, and ocular disorders comprises a kininogen domain
  high mol. Wt. Kiningen (HK) domain 3 (see AAY95426). The D3 peptide inhibits endothelial cell proliferation and thus possesses anti-anglogenic activity. It is an example of D3 peptides of the invention (see AAY95465-66) that are analogues of certain sites in the HK domain 3, in this case amino acid residues Asn275-Lys282. The peptides inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting anglogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell proliferation of the retina are treated.
   Anti-angiogenic; angiogenesis; inhibitor; kininogen; homologue; endothelial cell proliferation; apoptosis; cancer; ocular disorder; rheumatoid arthritis; cytostatic; antiarthritic; antirheumatic; therapy; human; D3 peptide.
   The present sequence is that of a D3 peptide derived from human
   100.0%; Score 45; DB 21; Length 32; 100.0%; Pred. No. 0.2; tive 0; Mismatches 0; Indels
   Score 45; DB 21; Length 16; Pred. No. 0.099;
  0; Indels
  Mismatches
  AAY95408 standard; Peptide; 32 AA.
  Claim 4; Page 26; 44pp; English.
  ;
0
   100.0%;
100.0%;
   99WO-US28465
  98US-0112427.
  Anti-anglogenic D3 peptide.
   25-SEP-2000 (first entry)
  Ouery Match
Best Local Similarity 100.0
  Conservative
   Ouery Match
Best Local Similarity
'-hea 8; Conserve
   WPI; 2000-442247/38.
   (UTEM ) UNIV TEMPLE
  (MCCR/) MCCRAE R K.
  32 AA;
                                      16 AA;
   1 NNATFYFK 8
  1 NNATEYEK 8
   1 NNATEYEK 8
  WO200035407-A2.
     proliferation.
   Homo sapiens.
   02-DEC-1999;
  16-DEC-1998;
   22-JUN-2000.
  Accrae RK;
  AAY95408;
  Sequence
  Sequence
   3 analog
  RESULT 5
   AAY95408
  ò
     SXS
  å
   qq
   ö
   The present sequence is that of a D3 peptide derived from human this mol.wt. Kninogen (HK) domain 3 (see AAY95426). The D3 peptide inhibits endothelial cell proliferation and thus possesses anti-anglogenic activity. It is an example of D3 peptides of the invention (see AAY95405-26) that are analogues of certain sites in the HK domain 3, in this case amino acid residues Asa755-Lys282.

The peptides inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting andothelial cell proliferation, and inducing endothelial cell proliferation of inhibition of fibroblast growth factor-induced HUVEC cell
  Composition for inhibiting anglogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rhewmatoid arthritis, and ocular disorders comprises a kininogen domain
induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting angiogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell apoptosis. Cancer, rheumatoid arthritis, and ocular disorders characterized by undesired vascularization of the retina are treated. The ICSO value for the present peptide was less than 0.8 um for inhibition of fibroblast growth factor-induced HUVEC cell
   Gaps
  Anti-anglogenic; anglogenesis; inhibitor; kininogen; homologue; endochelial cell proliferation; apoptosis; cancer; ocular disorder; theumatoid arthritis; cytostatic; antlarthritic; antirheumatic; therapy; human; D3 peptide.
   ó
  100.0%; Score 45; DB 21; Length 16; 100.0%; Pred. No. 0.099; 1ve 0; Mismatches 0; Indels
   AAY95410 standard; Peptide; 16 AA.
   Claim 7; Page 26; 44pp; English.
   98US-0112427.
  99WO-US28465
   Ant1-anglogenic D3 peptide.
  25-SEP-2000 (first entry)
   Best Local Similarity 100.
Matches 8; Conservative
   (UTEM ) UNIV TEMPLE. (MCCR/) MCCRAE R K.
   WPI; 2000-442247/38.
   16 AA;
  1 NNATEYEK 8
   WO200035407-A2.
  proliferation.
  Homo sapiens
   02-DEC-1999;
   16-DEC-1998;
  22-JUN-2000.
  AcCrae RK;
   Sequence
  AAY95410;
  3 analog
  Query Match
  RESULT 4
   8888888888888
  ò
   a
```

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Gaps

Homo sapiens

Pept1de Protein

AAR33350;

AAR33350 RESULT

a

13-AUG-1992; 13-AUG-1991;

WO9303748-A. 04-MAR-1993

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The present invention relates to inhibition of the enzymatic activity of legumain or a legumain related endopeptidase by cystatin. The inhibition involves an interaction between legumain and a papain-non-reactive site of cystatin. Legumain (EC 3.4.22.34) is a cysteine endopeptidase, and performs a protein-processing function. The present sequence is human kininogen D3, which was used in the present invention. Kininogen is a type 3 cystatin.
   Inhibition of mammalian legumain or legumain-related endopeptidase by cystatin involves interaction with second papain-non-reactive site of
   Anti-anglogenic; anglogenesis; inhibitor; kininogen; homologue; endothelial cell prolliferation; apoptosis; cancer; ocular disorder; rheumatoid arthritis; cyotatatic; antiarthritic; antirheumatic; therapy; human; D3 peptide.
   Enzyme; legumain; endopeptidase; cystatin; human; kininogen.
  Score 45; DB 21; Length 122; Pred. No. 0.8; 0; Mismatches 0; Indels
   Human high mol.wt. kininogen domain 3.
         AAB37447 standard; protein; 122 AA.
  AAY95426 standard; Peptide; 123 AA.
  Disclosure; Fig 4; 45pp; English.
  100.0%;
100.0%;
  20-APR-2000; 2000WO-GB01571.
  99GB-0009133.
   Barrett AJ;
  (first entry)
   Conservative
  (BABR-) BABRAHAM INST
  WPI; 2000-687316/67.
  Human kininogen D3
  Local Similarity
nes 8; Conserv
   Sequence 122 AA;
   1111111
35 NNATFYFK 42
  œ
  WO200064945-A1
  WO200035407-A2
  Abrahamson M,
   1 NNATFYFK
   Homo sapiens.
  22-APR-1999;
   21-FEB-2001
  Homo sapiens
  02-NOV-2000
  25-SEP-2000
  22-JUN-2000
  cystatin
  AAY95426;
  Query Match
   Best Loca
Matches
   AAY95426
   RESULT
         NAME OF COLOR OF STREET OF
   q
   ö
  The sequence given represents domain 3, amino acids 246-362, of the human kininogen heavy chain. Domain 3 was isolated from low molecular weight kininogen, derived from human plasma, by cleavage with trypsin. Domain 3 peptide inhibits platelet activation causing a marked decrease in the platelets ability to aggregate and secrete their granule contents. The granule contents comprise proteins which participate in hemostasis, thrombosis and the inflammatory response. Domain 3 also inhibits endothelial cell activation shown by a decrease in secretion of endothelial cell contents such as tissue plasminogen activator and von Willebrand factor. Domain 3 functions to inhibit cell activation by blocking thrombin binding to its target cells, the peptide is a selective inhibitor of thrombin-induced platlet
   Domain 3; human; kininogen; heavy chain; low molecular weight; plasma; trypsin; platelet; activation; granule contents; hemostasis; thrombin; tissue plasminogen activator; thrombosis; inflammatory response; endothelial cell; vow Willebrand factor;
  Gaps
  ö
   100.0%; Score 45; DB 14; Length 117; 100.0%; Pred. No. 0.77; 1ve 0; Mismatches 0; Indels (
   Use of Lrypsin-cleavage fragment of human kininogen - for increasing vascular bradykinin release, for lowering blood
   Domaine 3, bradykinin release activating peptide.
   /note= "Leader peptide"
19..117
   /note= "Mature protein"
  Location/Qualiflers
   AAR33350 standard; protein; 117 AA.
  pressure and treating hypertension
   Disclosure; Fig 1; 46pp; English.
  91US-0744545
  92WO-US06809
   01-JUL-1993 (first entry)
   Local Similarity 100.
   Jiang Y, Schmaier AB;
   (UTEM ) UNIV TEMPLE.
  WPI; 1993-093714/11.
   117 AA;
13 NNATFYFK 20
   30 NNATEYEK 37
   1 NNATFYFK 8
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Gaps

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99WO-US28465.

02-DEC-1999;

Query Match Best Local S Matches 8

ò a RESULT 7

Sequence

```
1 NNATFYFK 8
   WO200175067-A2
  Homo sapiens.
                    biodiversity
  11-OCT-2001.
   ABG21099;
  Matches
  RESULT 10
   ABG21099
  ð
   a
  ö
   Composition for inhibiting angiogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rhewmatoid arthritis, and ocular disorders comprises a kininogen domain
  Gaps
   The present sequence is that of domain 3 of human high mol.wt. kiningen (HK). The invention provides peptides (see AAY95405-24) that are analogues of certain sites in the HK domain 3, specifically Asn275-Lys282, Cys246-Cys249, Leu331-Tyr338 and Tyr299-Ser314. The peptides, in which native Cys residues may be replaced by Ala residues, inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting anglogenesis, inhibiting endothelial cell apoptosis. Cancer, rheumatoid arthritis, and coular disorders characterized by undesired vascularization of
  Human; chromosome mapping; gene mapping; gene therapy; forensic; food supplement; medical imaging; diagnostic; genetic disorder.
  New isolated polynucleotide and encoded polypeptides, useful in
  ö
   100.0%; Score 45; DB 21; Length 123; 100.0%; Pred. No. 0.81; 0; Mismatches 0; Indels (
  Novel human diagnostic protein #21093.
  ABG21102 standard; Protein; 248 AA.
  Oisclosure; Page 4; 44pp; English.
  Tang YT;
        98US-0112427.
   30-MAR-2001; 2001WO-US08631.
  31-MAR-2000; 2000US-0540217.
23-AUG-2000; 2000US-0649167.
  (first entry)
  the retina are treated.
                          (UTEM ) UNIV TEMPLE. (MCCR/) MCCRAE R K.
   WPI; 2001-639362/73.
   WPI; 2000-442247/38.
   Drmanac RT, Liu C,
   123 AA;
   (HYSE-) HYSEQ INC
  N-PSDB; AAS85289
  1 NNATFYFK 8
  WO200175067-A2.
   Homo sapiens.
        16-DEC-1998;
  18-FEB-2002
  11-0CT-2001.
   McCrae RK;
  Sequence
  ABG21102;
   3 analog
  RESULT 9
   ABG21102
à
   g
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The invention relates to isolated polynucleotide (I) and probes, polypeptide (II) sequences. (I) is useful as hybridisation probes, polymerase chain reaction (PCR) primers, oligomers, and for chromosome and gene mapping, and in recombinant production of (II). The polynucleotides are also used in diagnostics as expressed sequence tags for identifying expressed genes: (I) is useful in gene therapy techniques to restore normal activity of (II) or to treat disease states involving to restore normal activity of (II) or to treat disease states involving a polypeptide in tissue, as molecular weight markers and as a food supplement. (II) and its binding partners are useful in medical imaging of sites expressing (II). (I) and (II) are useful in medical inaging of sites expressing (II). (I) and (II) are useful in medical inaging of sites expressing (II). (I) and (II) are useful in medical closorders involving aberrant protein expression or biological activity. The polypeptide and polynucleotide sequences have applications in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other trails to assess biodiversity and to produce other types of data and products dependent on DNA and anno acid sequences. ABG00010-ABG30377 represent novel human order desquences data for this parent did not appear in the printed specification, but was obtained in electronic format directly from WIPO
   Gaps
diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
   diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
   Human; chromosome mapping; gene mapping; gene therapy; forensic; food supplement; medical imaging; diagnostic; genetic disorder.
  New isolated polynucleotide and encoded polypeptides, useful in
   ó
   0; Indels
   100.0%; Score 45; DB 22; 100.0%; Pred. No. 1.7;
   at ftp.wipo.int/pub/published_pct_sequences
   Mismatches
   Claim 20; SEQ ID No 51461; 103pp; English.
  Novel human diagnostic protein #21090.
  ABG21099 standard; Protein; 369 AA.
  ö
   Tang YT;
  30-MAR-2001; 2001WO-US08631.
   31-MAR-2000; 2000US-0540217.
23-AUG-2000; 2000US-0649167.
  18-FEB-2002 (first entry)
   Conservative
   WPI; 2001-639362/73.
   Drmanac RT, Liu C,
   Query Match
Best Local Similarity
  Sequence 248 AA;
  (HYSE-) HYSEQ INC
   N-PSDB; AAS85286.
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The present invention describes an isolated polypeptide (I) that

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of biological and which:

(a) inhibits angiogenesis; (b) binds to the D5 binding site on

condition of apoptosis in EC; and/or (d) inhibits the signalling

contraduction of apoptosis in EC; and/or (d) inhibits the signalling

contraduction of apoptosis in EC; and/or (d) inhibits the signalling

contraduction of apoptosis in EC; and/or (d) inhibits

contraduction ophthalmological, gynaecological, antiulcer, antidiabetic,

antiarthritic, antiangiogenic, antiangiogenic, antialogenic, antidiabetic,

antiarthritic, antiangiogenic, antiangiogenic, antidiabetic,

antiarthritic, antidiagiogenic, antidiogenic in a subject.

contraduction or angiogenesis in a subject. (I), a D5 fusion polypeptide

(II) or a dimeric or trimeric fusion polypeptide (III) can be used for

inhibiting EC migration, proliferation, invasion, or angiogenesis, or

confosition (X) comprising (I), (II), or (III), can be used for treating

confosition (X) comprising (I), (II), or (III), can be used for treating

confosition proliferation, invasion or angiogenesis. (I), (II), or (III)

con be used for isolating or enriching cells expressing D5 domain binding

sites from a cell mixture. The present sequence represents the mature

contraduction of the prosent invasion in the present invasion in the present invasion in the present invasion in the molecular weight kininogen (IK) protein, which is given in
                            conditions associated with endothelial cell migration, proliferation, invasion or angiogenesis, e.g. arthritis, macular degeneration, benign
     Novel human kininogen D5 domain polypeptides useful for treating
   Human; chromosome mapping; gene mapping; gene therapy; forensic; food supplement; medical imaging; diagnostic; genetic disorder.
  100.0%; Score 45; DB 23; Length 626; 100.0%; Pred. No. 4.3; ive 0; Mismatches 0; Indels (
   the exemplification of the present invention
   Novel human diagnostic protein #21092.
   ABG21101 standard; Protein; 644 AA.
  Disclosure; Page 13; 84pp; English.
  Tang YT;
  30-MAR-2001; 2001WO-US08631
   2000US-0540217
   2000US-0649167
  18-FEB-2002 (first entry)
   Ouery Match 100.
Best Local Similarity 100.
Matches 8; Conservative
   Drmanac RT, Liu C,
  WPI; 2001-639362/73
  626 AA;
  1111111
275 NNATEYEK 282
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  WO200175067-A2.
   Homo sapiens.
   23-AUG-2000;
   31-MAR-2000;
   11-OCT-2001.
   Sequence
   ABG21101;
  RESULT 12
   ABG21101
   ; oy
     g
  polymerase chain reaction (FOR) primers, oligomers, and for chromosome mode mapping, and in recombinant production of (II). The mode mapping, and in recombinant production of (II). The polymerase chain reaction (FOR) primers, oligomers, and for chromosome polymucleotides are also used in diagnostics as expressed sequence tags for identifying expressed genes. (I) is useful in gene therapy techniques to restore normal activity of (II) or to treat disease states involving (II). (II) is useful for generating antibodies against it, detecting or quantitating a polypeptide in tissue, as molecular weight markers and as a food supplement. (II) and its binding partners are useful in medical magning of sites expressing (II). (I) and (II) are useful for treating clisorders involving aberrant protein expression or biological activity. The polypeptide and polyucieched sequences have applications in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits to assess blodiversity and can do produce other types of data and products dependent on DNA and amino acid sequences. ABGO0010-ABG30377 represent novel human Note: The sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from NIPO at the vipo.int/pub/published_pct_sequences.
  ö
   Human; kininogen; high molecular weight kininogen; HK; D5 domain; D5 receptor; angiogenesis; endothelial cell; cytostatic; antitumour; antiatherosclerotic; vasotropic; vulnerary; tranquilliser; thrombolytic; ophthalmological; gypaecological; antiulcer; antidiabetic; antiarthritic; antiangiogenic; antiapoptotic; endocrine; apoptosis; gene therapy.
   Human high molecular weight kininogen (HK) mature protein SEQ ID NO:1.
  Gaps
  ö
   100.0%; Score 45; DB 22; Length 369; 100.0%; Pred. No. 2.5; ive 0; Mismatches 0; Indels
   invention relates to isolated polynucleotide (I) and
  Claim 20; SEQ ID No 51458; 103pp; English.
   Location/Qualifiers
   ABB78707 standard; Protein; 626 AA.
  384..508
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  24-JUL-2000; 2000US-220194P.
   24-JUL-2001; 2001WO-US23185
   (first entry)
   8; Conservative
  Mazar AP, Juarez JC;
  (ATTE-) ATTENUON LLC.
   WPI; 2002-393611/42.
  Local Similarity
   369 AA;
  333 NNATFYFK 340
   1 NNATFYFK 8
  WO200214369-A2
   18-JUL-2002
biodiversity
  Homo sapiens
   21-FEB-2002
  Seguence
   ABB78707;
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Domain

Query Match

Matches

RESULT 11 ABB78707

a

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Gaps

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RESULT 14
   AAW54336
     ò
  The invention relates to isolated polynucleotide (I) and polypeptide (II) sequences. (I) is useful as hybridisation probes, polymerase chain reaction (FGR) primers, oligomers, and for chromosome, and gene mapping, and in recombinant production of (II). The polynucleotides are also used in diagnostics as expressed sequence tags for identifying expressed genes. (I) is useful in gene therapy techniques to restore normal activity of (II) or to treat disease states involving to restore normal activity of (II) or to treat disease states involving appendent. (II) and its sub-diding partners are useful in medical maging of sites expressing (II). (I) and (II) are useful in medical activity. (II) and its binding partners are useful in medical maging of sites expressing (II). (I) and (II) are useful for treating clasorders involving aberrant protein expression or biological activity. The polypeptide and polynuclectide sequences have applications in cappositics, forensics, gene mapping, identification of mutations and to produce other types of data and products dependent on DNA and and canno acid sequences of the invention.

Comino acid sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from WIPO capporary in the printed capporary in the prin
  ö
  Human, kininogen, high molecular weight kininogen; HK; D5 domain, D5 receptor; angiogenesis; endothelial cell; cytostatic; antitumour; antiatherosclerotic; vasotropic; vulnerary; tranquilliser; thrombolytic; ophthalmological; gynaecological; antiulcer; antidiabetic; antiatrhritic; antianglogenic; antiapoptotic; endocrine; apoptosis; gene therapy.
  Gaps
  label- mature_human_high_molecular_weight_kininogen
New isolated polynucleotide and encoded polypeptides, useful in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
  ö
  / Match 100.0%; Score 45; DB 22; Length 644; Local Similarity 100.0%; Pred. No. 4.5; nes 8; Conservative 0; Mismatches 0; Indels (
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Jomain
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  ABB78710
  à
   g
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The present invention describes an isolated polypeptide (1) that

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of mortivative, and which:

(a) inhibits angiogenesis; (b) binds to the D5 binding site on

condothelial cells (EC); (c) activates signalling pathways leading to the

introduction of apoptosis in EC; and/or (d) inhibits the signalling

pathway required for maintenance of EC viability. (1) has cytostatic,

antitumour, antiatherosclerotic, vasotropic, vulnerary, tranquilliser,

trombolyytic, ophthalmological, gynaecological, antilicer, antidiabetic,

antiarthritic, antiangiogenic, antiapoptotic and endocrine activities.

An antibody (IX) specific for an epitope of (1) is useful for inhibiting

controlling EC migration, proliferation, invasion, or angiogenesis, or

composition (X) comprising (1), (11), con planding paramaceutical

composition (X) comprising (1), (11), con (111), can be used for treating

a subject having a disease or condition associated with undesired EC

migration, proliferation, invasion or angiogenesis. (1), (11), or (111),

can be used for isolating or enriching cells expressing D5 domain binding

sites from a cell mixture. The present sequence represents the human

control of the present invariet in the

energy the composition of the present invariety.
  Inhibition; thrombin-induced platelet; prevention; platelet aggregation; ADP-induced activation.
   Novel human kininogen D5 domain polypeptides useful for treating conditions associated with endothelial cell migration, proliferation, invasion or anglogenesis, e.g. arthritis, macular degeneration, benign
   Gaps
   ó
   100.0%; Score 45; DB 23; Length 644; 100.0%; Pred. No. 4.5; tive 0; Mismatches 0; Indels 0
  exemplification of the present invention.
   Disclosure; Fig 1B-E; 84pp; English.
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  Bradykinin analogous peptide 19.
  24-JUL-2000; 2000US-220194P.
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  Mazar AP, Juarez JC;
  (ATTE-) ATTENUON LLC
   WPI; 2002-393611/42.
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Best Local Similarity
Matches 8; Conserv
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   1 NNATFYFK 8
   WO200214369-A2
  30-JUL-1998
   21-FEB-2002
  Sequence
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   AAW54336;
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Search completed: March 13, 2003, 19:12:02 Job time : 22 secs
   Sequence
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  g
   ó
   Human; cytokine; cell proliferation; cell differentiation; gene therapy; vacchie; peptide therapy; stem cell growth factor; haematopoiesis; tissue growth factor: immunomodulatory; cancer; leukaemia; nervous system disorders; arthritis; inflammation.
   disclosed peptides and multimers can be used for the inhibition of thrombin-induced platelets or other cells. They can also be used for preventing platelet aggregation, or inhibiting ADP-induced activation. This is useful to prevent arterial occlusions arising from coronary thrombosis and stroke.
  Administration of a peptide or multimer related to bradykinin or other
   Inhibition of platelet activation and aggregation - by admin. of new or known bradykinin analogues
   Gaps
   Isolated nucleic acids and polypeptides, useful for preventing diagnosing and treating e.g. leukaemia, inflammation and immune disorders -
   ö
  Score 39; DB 18; Length 26;
Pred. No. 2;
0; Mismatches 0; Indels
   Disclosure; Page 44; 73pp; English.
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100.0%; Pre
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  Tang YT, Liu C, Drmanac RT;
  96WO-US09940
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18-MAY-2000; 2000US-0577409.
  26-FEB-2001; 2001WO-US04927
  06-NOV-2001 (first entry)
   Query Match
Best Local Similarity luv...
   Hasan AAK, Schmaier AH;
  (UNMI ) UNIV MICHIGAN.
  WPI; 1997-065304/06.
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   (HYSE-) HYSEQ INC
   26 AA;
  N-PSDB; AAI85011.
  2 NATFYFK 8
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  Homo sapiens.
WO9641640-A1
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                    27-DEC-1996
   07-SEP-2001,
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  RESULT 15
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   q
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  The invention relates to human polynucleotides (AAI79941-AAI93841) and the encoded proteins (AAO00010-AAO13910) that exhibit activity elating to cytokine, cell proliferation or cell differentiation or which may induce production of other cytokines in other cell populations. The polynucleotides and polypeptides are useful in gene therapy, vaccines or peptide therapy. The polypeptides have various cytokine-like activities, e.g. stem cell growth factor activity, haematopoiess regulating activity, tissue growth factor activity, immunomodulatory activity and activity and may be useful in the diagnosis and/or inflammation.
  Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/published_pct_sequences.
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Sequence 1 Sequence 1 Sequence 1

Sequence Sequence

Sequence Sequence

Sequence 3 Sequence 8 Sequence 9 Sequence 1

Sequence Sequence

Sequence Sequence

us-09-461-061a-1.rai

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APPLICANT: Schmaler, Alvin H.
APPLICANT: Schmaler, Alvin H.
APPLICANT: Jiang, Yongping
TITLE OF INVENTION: Modulation of Blood
TITLE OF INVENTION: Pressure and Inhibition of Platelet Activation
TITLE OF INVENTION: with Kininogen Fragment
NUMBER OF SEQUENCES: 2
ADDRESSE: Seidel, Gonda, Lavorgna &
ADDRESSE: Monaco, P.C.
STREET: 1800 Two Penn Center Plaza
CITY: Philadelphia
STATE: Pennsylvania
   100.0%; Score 45; DB 1; Length 117; 100.0%; Pred. No. 0.18;
   ZIP: 1910
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COMPUTER: IBM PS/2
COMPUTER: WordPerfect 5.1
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/193,114B
CLASSIFICATION NUMBER: US. Application
APPLICATION NUMBER: Serial No. 5472945 07/744,545
FILING DATE: 13 August 1991
ATTORNEY/AGENT INFORMATION:
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US-08-446-100-4
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   6056-137 CI1
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   NAME: MODICO, DELLE A.
REGISTRATION NUMBER: 30,480
REPERENCE/DOCKET NUMBER: 6056
TELECOMMUNICATION INFORMATION:
TELEFONE: (215) 568-5549
TELEX: NO. 54729456
INFORMATION FOR SEQ ID NO: 1:
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LENGTH: 117 amino acids
TTPE: peptide
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: U.S.A.
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Best Local Similarity
Matches 8; Conserv
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   US-08-193-114B-1
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Sequence 5
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US-08-149-727-2

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US-09-149-727-4

US-08-483-140-30

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US-08-418-905-8

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US-08-483-232-8

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US-09-20-041-8

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Gaps

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Indels

Gaps

; 0

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GENERAL INFORMATION:
APPLICANT: Jefferson, Richard A.
APPLICANT: Jefferson, Richard A.
APPLICANT: Jefferson, Richard A.
APPLICANT: Hilan, Andrzej
APPLICANT: Keese, Paul Konrad
TITLE OF INVENTION: USES THEREOF
TITLE OF INVENTION: USES THEREOF
TITLE REFERENCE: 190106.405
CURRENT APPLICATION NUMBER: US/09/149,727
CURRENT FILING DATE: 1998-09-08
EARLIER FILING: DATE: 1997-09-09
NUMBER OF SEQ ID NOS: 71
SOFTWARE: Patentin Ver. 2.0
SEQ ID NO 2.
LENGTH: 602
  APPLICANT: Jefferson, Richard A.
APPLICANT: Jefferson, Richard A.
APPLICANT: Milan, Andrej
APPLICANT: Kilian, Andrej
APPLICANT: Kilian, Andrej
APPLICANT: Keese, Paul Konrad
TITLE OF INVENTION: MICROBIAL BETA-GLUCURONIDASE GENES, GENE PRODUCTS AND
TITLE OF INVENTION: WISES THEREOF
FILE REPRENDE: 199106-408
CURRENT APPLICATION NUMBER: US 60/058,263
EARLIER APPLICATION NUMBER: US 60/058,263
EARLIER PILING DATE: 1997-09-09
NUMBER OF SEQ ID NOS: 71
SOFTWARE: PALENTIN Ver. 2.0
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EARLIER FILING DATE: 1995-06-09
EARLIER APPLICATION NUMBER: PCT/US96/09940
EARLIER FILING DATE: 1996-06-07
NUMBER OF SEQ ID NOS: 24
SOFTWARE: Patentin Ver. 2.1
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LENGTH: 26
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100.0%; Pred. No.
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Patent No. 6391547
  US-09-149-727-8; Sequence 8, Application US/09149727; Sequence 8, Application US/09149727; Defent No. 6391547; GENERAL INFORMATION:
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  FEATURE:
  ò
  g
  ò
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  ö
  Sequence 15, Application US/08676242C
Batter No. 6443719
GENERAL INFORATION
APPLICANT: The Regents of the University of Michigan
APPLICANT: Schmater, Alvin H.
APPLICANT: Hasan, Ahmad A.K.
TITLE OF INVENTION: Bradykinin Analogs As Selective Thrombin Inhibitors
FILE REFERENCE: 8820-2 US
CURRENT APPLICATION UNDER: US/08/676,242C
CURRENT FILING DATE: 2000-07-16
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  APPLICANT: Schmaier, Alvin H.
APPLICANT: Jiang, Yongping
TITLE OF INVENTION: Modulation of Blood
TITLE OF INVENTION: Pressure by Altering Bradykinin Levels
NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
ADDRESSEE: Temple University - Of the
ADDRESSEE: Commonwealth System of Higher Education
STREET: 406 University Services
STREET: Building
  100.0%; Score 45; DB 5; Length 117; 100.0%; Pred. No. 0.18; ive 0; Mismatches 0; Indels
   COUNTRY:

2IP: 1912.

COMPUTER READABLE FORM:

MEDIUM TYPE: Diskette, 3.50 inch, 720 Kb
COMPUTER: IBM PS/2

COMPUTER: IBM PS/2

COMPUTER: WordPerfect 5.1

CURRENT APPLICATION DATA:

; APPLICATION UNMBER: PCT/US92/06809

FILING DATE: 19910813
   PRIOR APPLICATION DATA:
APPLICATION NUMBER: U.S. Application
APPLICATION NUMBER: Serial No. 744,545
FILING DATE: 13 AUGUST 1991
ATTORNEY/AGENT INFORMATION:
   NAME: Monaco, Daniel A. REGISTRATION WOMBER: 30,480
REFERENCE/DOCKET NUMBER: 6056-137
TELECOMMUNICATION INFORMATION:
  Sequence 1, Application PC/TUS9206809 GENERAL INFORMATION:
  INFORMATION FOR SEQ ID NO: 1:
   : 117 amino acids
AMINO ACID
   TELEPHONE: (215) 568-83:
TELEFAX: (215) 568-5549
  Query Match
Best Local Similarity 100...
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Gaps

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   qq
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| GENERAL INFORMATION:
| APPLICANT: Jefferson, Richard A. |
| APPLICANT: Meferson, Richard A. |
| APPLICANT: Messe, Paul Konrad |
| TITLE OF INVENTION: MICROBIAL BETA-GLUCURONIDASE GENES, GENE PRODUCTS AND |
| TITLE OF INVENTION: MICROBIAL BETA-GLUCURONIDASE GENES, GENE PRODUCTS AND |
| TITLE OF INVENTION: MICROBIAL BETA-GLUCURONIDASE GENES, GENE PRODUCTS AND |
| TITLE OF INVENTION: MUMBER: US/09/149,727 |
| CURRENT APPLICATION NUMBER: US/09/08 |
| EARLIER APPLICATION NUMBER: US/09/09 |
| FARLIER FILING DATE: 1997-09-09 |
| SOFTWARE: PatentIn Ver. 2.0 |
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  Sequence 30, Application US/08483140
Fatent No. 5688403
GENERAL INFORMATION:
TITLE OF INVENTION: Platelet-Activating Factor Acetyl
TITLE OF INVENTION: Hydrolase
NUMBER OF SEQUENCES:
CORRESPONDENCE ADDRESS:
ADDRESSE: Marshall, O'Toole, Gertein, Murray & Borun
STREET: 6300 Sears Tower, 233 South Wacker Drive
CITY: Chicago
STATE: Illinois
COUNTRY: USA
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0; Mismatches 2; Indels
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PRICE APPLICATION NAMES: 6.0-07: 1994

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   0; Indels
  2: Marshall, O'Toole, Gertein, Murray & Borun 6300 Sears Tower, 233 South Wacker Drive
   SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/470,187
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Pred. No. 1e+02;
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PRIOR APPLICATION DATA:
APPLICATION NUMBER:
FILING DATE:
ATTORNEY/AGENT INFORMATION:
NAME: NO. 5532152and, Greta E.
REGISTRATION NUMBER: 35,302
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COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
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  68.98;
71.48;
  Query Match 68.9%;
Best Local Similarity 71.4%;
Matches 5; Conservative
   TELEPHONE: (312) 474-6300
TELEFAX: (312) 474-0448
TELEX: 25-3658
INFORMATION FOR SEQ ID NO: 8:
  : 193 amino acids
amino acid
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amino acid
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  CITY: Chicago
STATE: Illinois
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  ; MOLECULE TYPE: protein US-08-485-938A-36
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US-08-470-187-8
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                                       GENERAL INFORMATION:
APPLICANT: Cousens, Lawrence S.
APPLICANT: Eberhardt, Christine D.
APPLICANT: Eberhardt, Christine D.
APPLICANT: Totalkar, Larry W.
APPLICANT: Tjoelker, Larry W.
APPLICANT: Wilder, Cherry L.
TITLE OF INVENTION: Platelet-Activating Factor Acetyl
TITLE OF INVENTION: Hydrolase
NUMBER OF SEQUENCES: 26
CORRESPONDENCE ADDRESS:
   Length 441;
   0; Indels
  E: Marshall, O'Toole, Gertein, Murray & 36300 Sears Tower, 233 South Wacker Drive
  SOFTWARE: Patentin Release #1.0, Version #1.25 CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/318,905
FILLIG DATE:
CLASSIFICATION: 424
RICH APPLICATION A74:
APPLICATION NUMBER: US 08/133,803
FILLIO DATE: G-OCT-1993
ATTORNEY/AGENT INCOMATION:
   APPLICANT: COUSENS, Lawrence S.
APPLICANT: Eberhardt, Christine D.
APPLICANT: Gray, Partick W.
APPLICANT: Tjoelker, Larry W.
APPLICANT: Tjoelker, Larry W.
APPLICANT: Wilder, Cheryl L.
TILE OF INVENTION: Platelet-Activating Factor
TITLE OF INVENTION: Acetylhydrolase
  Score 31; DB 1;
Pred. No. 2.3e+02;
   Mismatches
   NAME: NO. 5641669and, Greta E. REGISTRATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 32205
  Sequence 8, Application US/08483232
Patent No. 5656431
Sequence 8, Application US/08318905
  COMPUTER READABLE FORM:
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  : (312) 474-6300
(312) 474-0448
   TELEFAX: (312) 474-0448
TELEX: 25-5658
INFORMATION FOR SEQ ID NO: 8:
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LENGTH: 441 amino acids
TYPE: amino acid
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; MOLECULE TYPE: protein
US-08-318-905-8
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STATE: Illinois
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185 SATYYFK 191
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Pred. No. 2.3e+02;
2; Mismatches 0; Indels
  SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/485,938A
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PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/318,905
FILING DATE: 06-0CT-1994
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/133,803
FILING DATE: 06-0CT-1993
ATTORNEY,AGENT INFORMATION:
NAME: No. 5847088and, Greta E.
REGISTRATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 27866/32792
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
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FILING DATE: 6-0CT-1994
PRIOR APPLICATION DATE: G-0CT-1993
APPLICATION NUMBER: US 08/133,803
FILING DATE: 6-0CT-1993
ATTORNEY/AGENT INFORMATION:
NAME: No. 5698403and, Greta E.
REGISTRATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 32781
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
TELEFAX: (312) 474-6300
TELEFAX: (312) 474-6448
ITELERX: 25-3658
INFORMATION FOR SEQ ID NO: 8:
SEQUENCE CHARACTERISTICS:
LENGTH: 441 amino acids
TYPE: amino acids
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  US-08-485-938A-8
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Pred. No. 2.3e+02;
2; Mismatches 0; Indels
NUMBER OF SEQUENCES: 30
CORRESPONDENCE ADDRESS:
ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & Borun
STREET: 6300 Sears Tower, 233 South Wacker Drive
   Sequence 8, Application US/08483140

Sequence 8, Application US/08483140

Patent No. 5698403

GENERAL INFORMATION:

APPLICANT: ICOS Corporation

TITLE OF INVENTION: Platelet-Activating Factor Acetyl

TITLE OF INVENTION: Hydrolase

NUMBER OF SEQUENCES: 30

CORRESPONDENCE ADDRESS:

ADDRESSEE: MAISHALL, O'Toole, Gertein, Murray & Borun

STREET: 6300 Sears Tower, 233 South Wacker Drive
  COMPUTER READABLE FORM:
MEDIUM TYPE: FOOPPY disk
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
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COMPUTER: PatentIn Release #1.0, Version #1.25
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/483,232
FILING DATE: 06-OCT-1994
PRIOR APPLICATION NUMBER: US 08/318,905
FILING DATE: 06-OCT-1993
APPLICATION NUMBER: US 08/133,803
FILING DATE: 06-OCT-1993
ATTORNEY/AGENT INFORMATION:
APPLICATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 35,302
TELEFAN: 1312) 474-6448
TELECOMMUNICATION INFORMATION:
TELECOMMUNICATION INFORMATION:
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EQUIRACE CHARACTERISTICS:
LENGTH: 441 amino acids
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COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
COMPUTER: IBM PC COMPATIBLE
OPERATING SYSTEM: PC-DOS/MS-DOS
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COUNTRY: United States of America
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ZIP: 6060-6402
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Matches 5; Conservative
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  FILING DATE:
CLASSIFICATION: 435
   CITY: Chicago
STATE: Illinois
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185 SATYYFK 191
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   GENERAL INFORMATION:
APPLICANT: COUSEN, LAWTENCE S.
APPLICANT: COUSEN, LAITON, W.
APPLICANT: Eberhardt, Christine D.
APPLICANT: Lairon, Hai
APPLICANT: Lairon, Hai
APPLICANT: Tjoelker, Lairy W.
APPLICANT: Wilder, Cheryl L.
TITLE OF INVENTION: Platelet-Activating Factor
TITLE OF INVENTION: Acetylhydrolase
NUMBER OF SEQUENCES: 30
CORRESPONDENCES: 30
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COUNTRY: United States of America
ADDRESSE: READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US 08/483,232
FILING DATE: 06-OCT-1994
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/318,905
FILING DATE: 06-OCT-1993
ATTORNEY/AGRIT INFORMATION:
NAME: PAT-LATION NUMBER: US 08-133,803
FILING DATE: 06-OCT-1993
ATTORNEY/AGRIT INFORMATION:
NAME: PAT-LATION NUMBER: US 08-133,803
FILING DATE: 06-OCT-1993
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FILING DATE: 06-OCT-1993
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NAME: PAT-LATION NUMBER: US 08-133,803
FILING DATE: 06-OCT-1993
ATTORNEY/AGRIT INFORMATION:
NAME: PAT-LATION NUMBER: US 08-134,605
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Pred. No. 2.3e+02;
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  Sequence 8, Application US/09328474
Patent No. 6045794
GENERAL INFORMATION:
   TELEPHONE: (312) 474-6300
TELEFAX: (312) 474-648
TELEX: 25-3658
INFORMATION FOR SEQ ID NO: 8:
SEQUENCE CHARACTERISTICS:
LENGTH: 441 amino acids
TYPE: amino acids
   NAME: Rin-Laures, Li-Hsien
REGISTRATION NUMBER: 33.547
REFERENCE/DOCKET NUMBER: 27
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2; Mismatches 0; Indels
   E: Marshall, O'Toole, Gerstein, Murray & Borun 6300 Sears Tower, 233 South Wacker Drive
   STATE: 6300 Sears Tower, 233 South Wacker Drive CITY: Chicago STATE: 111inois COUNTRY: United States of America ZIP: 6606-6402 ZOMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk COMPUTER: IBM PC compatible COMPUTER: IBM PC compatible SOFTWARE: PATENTING SYSTEM: PC-DOS/MS-DOS SOFTWARE: PATENTIN Release #1.0, Version #1.25 CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/910,041
   Sequence 8, Application US/08910041

Patent No. 5977308

GENERAL INFORMATION:
APPLICANT: Cousens, Lawrence S.
APPLICANT: Eberhardt, Christine D.
APPLICANT: Le Trong, Hal.
APPLICANT: Le Trong, Hal.
APPLICANT: Wilder, Cheryl L.
TITLE OF INVENTION: Platelet-Activating Factor:
TITLE OF INVENTION: Acetylhydrolase
NUMBER OF SEQUENCES: 30
CORRESPONDENCE ADDRESS:
  FILING DATE:
CLASSIFICATION: 424
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US 08/483,232
FILING DATE: 07-JUN-1995
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APPLICATION NUMBER: US 08/318,905
FILING DATE: 06-OCT-1994
PRIOR APPLICATION NUMBER: US 08/133,803
FILING DATE: 06-OCT-1993
APPLICATION NUMBER: US 08/133,803
FILING DATE: 06-OCT-1993
ATTORNEY/AGENT INFORMATION:
   NAME: Rin-Laures, Li-Hsien
RECISTRATION NUMBER: 33,547
REFERENCE/DOCKET NUMBER: 27866/34026
TELECOMMUNICATION INFORMATION:
TELEFAX: (312) 474-0448
TELEX: 25-3658
INFORMATION FOR SEQ ID NO: 8:
SEQUENCE CHARACTERISTICS:
LENGTH: 441 amino acids
TYPE: amino acid
  TELEPHONE: (312) 474-6300
TELEFAX: (312) 474-6448
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LENGTH: 441 amino acids
TYPE: amino acids
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   TOPOLOGY: linear

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APPLICANT: Rosen et al.

TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies FILE REFERENCE: PCOOB.

CURRENT APPLICATION NUMBER: US/09/764,847

CURRENT FILING DATE: 2001-01-17

Prior application data removed - consult PALM or file wrappe NUMBER OF SEQ ID NOS: 2003

SOFTWARE: Patentin Ver. 2.0

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Milne, Todd
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Royer, John
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Patent No. US/20020128250A1
GENERAL INFORMATION:
APPLICANT: Busby, Robert
APPLICANT: Call, Brian
APPLICANT: Hecht, Peter
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US-09-768-2359-6

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Gaps

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Indels

Result Š. 

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ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & Borun
STREET: 6300 Sears Tower, 233 South Wacker Drive
CITY: Chicago
   APPLICANT: Trawick, John D.
APPLICANT: Carr, Grant J.
APPLICANT: Yamanoto, Robert T.
APPLICANT: Xu, H. Howard
TITLE OF INVENTION: Identification of Essential Genes in FILE PREPERSENT.
  COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
   Trong, Hai
Trong, Hai
Trolker, Larry W.
Wilder, Cheryl L.
TITLE OF INVENTION: Platelet-Activating Factor
Acetylhydrolase
   Score 31; DB 10;
Pred. No. 88;
1; Mismatches 1
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MEDIUM TYPE: Floppy disk
  FILE REFERENCE: ELITRA.011A
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CURRENT FILING DATE: 2001-03-21
  SOFTWARE: FastSEQ for Windows Version 4.0 SEQ ID NO 13997
LENGTH: 224
   Sequence 8, Application US/09729402
Patent No. US20010021379A1
GENERAL INFORMATION:
APPLICANT: Cousens, Lawrence S.
Eberhardt, Christine D.
Gray, Patrick W.
  CURRENT FILING DATE: 2001-03-21
PRIOR APPLICATION NUMBER: 60/191,078
PRIOR FILING DATE: 2000-03-21
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PRIOR SPLICATION NUMBER: 60/206,848
PRIOR PELLOR DATE: 2000-05-28
PRIOR PRILING DATE: 2000-05-26
PRIOR APPLICATION NUMBER: 60/207,727
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PRIOR PELLING DATE: 2000-10-23
PRIOR PELLOR DATE: 2000-11-27
PRIOR PAPLICATION NUMBER: 60/253,625
PRIOR PELLOR DATE: 2000-11-27
PRIOR APPLICATION NUMBER: 60/257,931
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PRIOR PELLOR DATE: 2001-02-16
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  Ohlsen, Kari L.
Zyskind, Judith W.
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  .US-09-815-242-13997
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  TYPE: PRT
ORGANISM:
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   APPLICANT: Summers, Eric
TITLE OF INVENTION: Methods for Improving Secondary Metabolite Production in Fungi
FILE REFERENCE: 109272.147
CURRENT APPLICATION NUMBER: US/09/801,368
CURRENT FILING DATE: 2001-03-07
PRIOR APPLICATION NUMBER: US 09/487,558
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PRIOR APPLICATION NUMBER: US 00/160,587
PRIOR PLING DATE: 1999-10-20
NUMBER OF SED ID NOS: 440
SOFTWARE: Patentin version 3.0
  APPLICANT: CITPUS, Petra
APPLICANT: CITPUS, Petra
APPLICANT: Lerchl, Jens
APPLICANT: Meski, Malf
TITLE OF INVENTION: involved in the regulation of cell division, growth and
TITLE OF INVENTION: biomass formation in plants
FILE REFERENCE: 0093/000009
CURRENT APPLICATION NUMBER: US/09/768,235B
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PRIOR APPLICATION NUMBER: PCT/PED00/00675
PRIOR FILING DATE: 2000-01-28
NUMBER OF SEQ ID NOS: 45
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Schmidt, Ralf-Michael
Freund, Annette
   Ehrhardt, Thomas
Bischoff, Friedrich
Renz, Andreas
Duwenig, Elke
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                    Sherman, Am
Siľva, Jeff
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APPLICANT: Adler, Charles S.
APPLICANT: Adler, Jon Elliot
APPLICANT: Ryba, Nick
APPLICANT: Mueller, Ken
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Wark
APPLICANT: Hoon, Wark
APPLICANT: Hoon, Wark
APPLICANT: Hoon, Wark
APPLICANT: No. US20030022278alel Family of Taste Receptors
FILE REFERENCE: 02307E-098010US
CURRENT APPLICATION NUMBER: US/09/510,332
CURRENT FILING DATE: 2000-02-22
BRIONE APPLICATION NUMBER: US 09/393,634
PRIOR FILING DATE: 1999-09-10
  TITLE OF INVENTION: 123 Human Secreted Proteins
FILE REFERENCE: P2010P1
CURRENT APPLICATION NUMBER: US/09/983,802
CURRENT FILING DATE: 2001-10-25
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PRIOR PILING DATE: EARLIER FILING DATE: 1999-07-07
PRIOR PLING DATE: EARLIER FILING DATE: 1998-07-07
PRIOR APPLICATION NUMBER: EARLIER APPLICATION NUMBER: 60/051,926
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Pred. No. 51;
2; Mismatches (
  ; OTHER INFORMATION: mouse T2R28 (mGR28) US-09-510-332-158
  US-09-983-802-597; Application US/09883802; Sequence 597, Application US/09883802; Publication No. US20030022185A1; Publication No. US20030022185A1; APPLICANT: Fischer et al.
                               Application US/09510332
5. US20030022278A1
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                               Sequence 158, Applica
Publication No. US200
GENERAL INFORMATION:
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   ORGANISM: Mus sp.
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| SATEWE 68
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   TYPE: PRT
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   TITLE OF INVENTION: DROSOPHILA HOWOLGUES OF GENES AND PROTEINS IMPLICATED IN METABG
TITLE OF INVENTION: METHODS OF USE
FILE REPERENCE: EXOO-044C
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PRIOR PLICATION NUMBER: 60/172,482
PRIOR PLILNG DATE: 1999-12-17
PRIOR APPLICATION NUMBER: 60/172,482
PRIOR PLILNG DATE: 2000-01-27
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APPLICATION NUMBER: US 08/318,905
FILING DATE: 06-0CT-1994
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FILING DATE: 06-0CT-1993
ATORNEY/AGENT INFORMATION:
NAME: NO. US2010021379Aland, Greta E.
REGISTRATION NUMBER: 35,302
REFERENCE/DOCKET NUMBER: 27866/32793
TELECOMMUNICATION INFORMATION:
TELEFPHONE: (312) 474-6300
TELEFPHONE: (312) 474-6448
APPLICATION NUMBER: US/09/729,402
FILING DATE: 04-Dec-2000
CLASSIFICATION: CURNOWN>
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60 NATFFF 65
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   US-09-764-853-756
  SEQ ID NO 756
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GENERAL INFORMATION:
APPLICANT: Rank, David R.
APPLICANT: Hanzel, David K.
APPLICANT: Chen, Wensheng
TILLE OF INVENTION: HUMAN GENOME-DERIVED SINGLE EXON NUCLEIC ACID PROBES USEFUL FO
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   Length 127;
   2; Indels
  SOFTWARE: Annomax Sequence Listing Engine vers. 1.1 SEQ ID NO 35808
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PRIOR FILING DATE: 2000-02-04

PRIOR FILING DATE: 2000-05-26

PRIOR FILING DATE: 2000-06-36

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   1 NNATEYF 7
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US-09-864-761-35808
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GENERAL INFORMATION:
APPLICANT: Zuker, Charles S.
APPLICANT: Adder, Jon Elliot
APPLICANT: Ryba, Mick
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Wark
APPLICANT: The Regents of the University of California
TILLE OF INVENTION: T2R, a No. US20030022278Alel Family of Taste Receptors
FILE REFERENCE: 02307E-098010US
CURRENT APPLICATION NUMBER: US/09/510,332
CURRENT FILING DATE: 2000-02-22
PRIOR APPLICATION NUMBER: US 09/393,634
PRIOR FILING DATE: NO. US2005-09-10
   APPLICANT: Rene S. Hubert
APPLICANT: Pla M. Challita-Eld
APPLICANT: Pla M. Challita-Eld
APPLICANT: Pla M. Challita-Eld
APPLICANT: Alan Levin
APPLICANT: Alan Levin
APPLICANT: Steve Chappell Mitchell
APPLICANT: Steve Chappell Mitchell
APPLICANT: Aya Jakobovits
TITLE OF INVENTION: HIGHLY EXPRESSED IN PROSTATE CANCER
FILE REFERENCE: 129.50501
CURRENT APPLICATION NUMBER: US/09/780,053
CURRENT APPLICATION NUMBER: 60/181,261
PRIOR FILLING DATE: 2001-02-09
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93 NSATEWE 99
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93 NSATFWF 99
  1 NNATFYF 7
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US-09-780-053-5
   g
   g
  δ
  ò
   APPLICANT: Adler, Jon Elliot
APPLICANT: Adler, Jon Elliot
APPLICANT: Ryba, Nick
APPLICANT: Ryba, Nick
APPLICANT: Ryba, Nick
APPLICANT: Hoon, Mark
APPLICANT: Hoon, Mark
APPLICANT: The Regents of the University of California
TILLE REFERENCE: 02307E-098010US
CURRENT APPLICATION NUMBER: US/09/510,332
CURRENT APPLICATION NUMBER: US/09/510,332
FILE REFERENCE: 2000-02-22
PRIOR APPLICATION NUMBER: US 09/393,634
PRIOR FILING DATE: 1999-09-10
NUMBER OF SEQ ID NOS: 172
SOFTWARE: Patentin Ver: 2.1
SSETURENT PRIOR NUMBER: US 09/393,634
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   Gaps
   Gaps
  APPLICANT: ADLER, JON ELLIOT
TITLE OF INVENTION: TAR TASTE RECEPTORS AND GENES ENCODING SAME
FILE REFERENCE: 078003/0279152/RXT
CURRENT APPLICATION NUMBER: US/09/R25,882
CURRENT FILING DATE: 2001-04-05
PRIOR APPLICATION NUMBER: 60/195,532
PRIOR FILING DATE: 2000-11-13
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SOFTWARE: PATENTING PATE: 2000-11-13
SOFTWARE: PATENTING PATE: 2000-11-13
TYPE: PRIOR PATENTING PATE: 2000-11-13
TYPE: PRIOR PATENTING PATE: 2000-11-13
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OTHER INFORMATION: EXPRESSED IN LUNG, SIGNAL = 7.9

OTHER INFORMATION: EXPRESSED IN BRAIN, SIGNAL = 7.9

OTHER INFORMATION: EXPRESSED IN BT474, SIGNAL = 9.4

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93 NSATFWF 99
  1 NNATFYF 7
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US-09-825-882-18
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Gaps

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APPLICANT: Maxon, Mary
APPLICANT: Milne, Todd
APPLICANT: Milne, Todd
APPLICANT: No. US20020128250Alman, Thea
APPLICANT: No. US20020128250Alman, Thea
APPLICANT: Salama, Sofie
APPLICANT: Salama, Amir
APPLICANT: Summers, Eric
APPLICANT: Summers, Eric
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APPLICANT: Summers, Eric
APPLICANT: Summers, Exic
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CURRENT APPLICATION NUMBER: US/09/801,368
CURRENT FILING DATE: 2001-03-07
PRIOR FILING DATE: 2000-01-19
PRIOR FILING DATE: 1999-10-20
PRIOR FILING DATE: 1999-10-20
NUMBER: OF SEQ ID NOS: 440
SOFTWARE: PatentIn version 3.0
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; Sequence 336, Application US/09801368
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; GENERAL INPORMATION:
; APPLICANT: Busby, Robert
; APPLICANT: Call, Brian
...... Doug
   ; ORGANISM: Saccharomyces cerevisiae US-09-801-368-336
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US-09-780-053-5
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Madden, Kevin
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APPLICANT:
APPLICANT:
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| GenCore version 5.1.4_p5_4578<br>Copyright (c) 1993 - 2003 Compugen Ltd. | OM protein - protein search, using sw model | March 13, 2003, 19:10:37; Search time 9 Seconds (without alignments) 85.453 Million cell updates/sec |
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|                                                                          | OM protein .                                | Run on:                                                                                              |

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283224

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

PIR\_73:\*
1: pir1:\*
2: pir2:\*
3: pir3:\*
4: pir4:\* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

& Query Result

| Description | kininogen, LMW pre | ĕ      | n' u   | kininogen, HMW I p | hypothetical prote | telomerase reverse | hypothetical prote |        | α      | hypothetical prote | rhamnosyltransfera | NADH2 dehydrogenas | probable cytochrom | hypothetical prote | hypothetical prote | v      | hypothetical prote | hypothetical prote | hypothetical prote | S      | hypothetical prote | hypothetical prote | •      | secretory protein | invasion protein 1 |        | type III secretion | NADH2 dehydrogenas | hypothetical prote |  |
|-------------|--------------------|--------|--------|--------------------|--------------------|--------------------|--------------------|--------|--------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------|--------------------|--------------------|--------------------|--------|--------------------|--------------------|--------|-------------------|--------------------|--------|--------------------|--------------------|--------------------|--|
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| Match       | 100.0              | 100.0  |        |                    | 80.0               |                    | 7                  | 'n     |        | S.                 | δ.                 |                    | Š.                 | 'n                 | S.                 | S.     | 73.3               | Э.                 | ъ.                 | œ.     | œ.                 | ω.                 | ij     | ij                | 71.1               | Ξ.     | ÷                  | 71.1               | 71.1               |  |
| Score       | 45                 | 45     | 39     | 39                 | 36                 | 36                 | 32                 | 34     | 34     | 34                 | 34                 | 34                 | 34                 | 34                 | 34                 | 34     | 33                 | 33                 | 33                 | 33     | 33                 | 33                 | 32     | 32                | 32                 | 32     | 32                 | 32                 | 32                 |  |
| SO.         | 7                  | ~      | m      | 4                  | 'n                 | ø                  | 7                  | œ      | 6      | 10                 | 11                 | 12                 | 13                 | 14                 | 15                 | 16     | 17                 | 18                 | 19                 | 20     | 21                 | 22                 | 23     | 24                | 25                 | 56     | 27                 | 28                 | 59                 |  |

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| protein kinase GCN cytokinin-induced | hypothetical prote hypothetical prote | hypothetical prote<br>amidotransferase h | probable permease | hypothetical prote | U2 snRNP auxiliary | conserved hypothet | conserved hypothet | U2 small nuclear r | N-acety1-gamma-glu | N-acetyl-gamma-glu | N-acetyl-gamma-qlu | outer membrane pro |
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| OKBYN2<br>T10059                     | B71529<br>E90088                      | F84366<br>F84941                         | AF0604            | T34501             | A46179             | F69409             | 829078             | JC6125             | RDECEP             | G91239             | D86087             | A82299             |
| 7                                    | 77                                    | ~ ~                                      | 7                 | ~                  | N                  | ~                  | ~                  | ~                  | Н                  | ~                  | ~                  | 7                  |
| 1659                                 | 148                                   | 190                                      | 202               | 211                | 240                | 249                | 262                | 264                | 334                | 334                | 334                | 350                |
| 71.1                                 | 68.9<br>68.9                          | 68.9                                     | 6.89              | 68.8               | 68.9               | 68.8               | 68.8               | 68.8               | 68.8               | 68.8               | 68.8               | 68.8               |
| 32                                   | 31                                    | 31                                       | 31                | 31                 | 31                 | 31                 | 31                 | 31                 | 31                 | 31                 | 31                 | 31                 |
| 30<br>31                             | 3.2                                   | 9.64<br>4.62                             | 36                | 37                 | 38                 | 39                 | 40                 | 41                 | 42                 | 43                 | 44                 | 45                 |

## ALIGNMENTS

|   | KGHUL1 KGHUL1 KAILOLOGEN, LMW precursor [validated] - human Nillernate names: alpha-2-thiol proteinase inhibitor; preprokininogen N; Contains: bradykinin (kallidin); kininogen I; kininogen II; prokininogen C; Species: Homo sapiens (man) C; Date: 06-ul-1981 #sequence_revision 27-Nov-1985 #text_change 08-Dec-2000 C; Accession: A01280; B25276; A27900; A27699; A31905; A34030 R; Ohkubo, II; Kurzechi, K.; Takasawa, T.; Shiokawa, H.; Sasaki, M. Biochemistry 23, 5691-5697, 1984 |
|---|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|   | A/Fitle: Isolation of a human cDNA for alpha-2-thiol proteinase inhibitor and its ide A/Reference number: A90490; MUID:85122621; PMID:6441591 A/Accession: A01280 A/Molecule type: mRNA                                                                                                                                                                                                                                                                                                    |
| _ | A;residues: 1-42/ CORA<br>A;Cross-references: GB:K02566; NID:g177889; PIDN:AAA35497.1; PID:g177890<br>B;Takagaki, Y.; Kitamura, N.; Nakanishi, S.<br>J. Biol. Chem. 260. 8601-8609, 1985                                                                                                                                                                                                                                                                                                   |
|   | Arritle: Cloning and Sequence analysis of cDNAs for human high molecular weight and l<br>A;Reference number: A92544; MUID:85234582; PMID:2989293<br>A;Accession: B25276                                                                                                                                                                                                                                                                                                                    |
|   | A; Molecule type: mRNA A; Residues: 1-427 <tak> A; Residues: 1-427 <tak> A; Coss.references: GS:M11437; NID:g186751; PIDN:AAB59551.1; PID:g386853</tak></tak>                                                                                                                                                                                                                                                                                                                              |
|   | Kilottspeich, F.; Kellermann, J.; Henschen, A.; Kauth, G.; Mueller-Esterl, W. in Kinns IV, part A. Greenbaum, L.M., and Margolius, H.S., eds., pp.91-95, Plenum, N. A.Title: Amino acid sequence of the light chain of human low molecular mass Kininogen A:Reference number: A27900                                                                                                                                                                                                       |
|   | A,Accession: A27900<br>A,Molecule type: protein<br>A.Residues: 390-427 <pre>cLOT&gt;</pre>                                                                                                                                                                                                                                                                                                                                                                                                 |
| : | R.Mindrolu, T.; Carretero, O.A.; Proud, D.; Walz, D.; Scicli, A.G.<br>Biochem. Biophys. Res. Commun. 152, 519-526, 1988<br>A:Title: A new kinin moiety in human plasma kininogens.                                                                                                                                                                                                                                                                                                         |
|   | A; Reference number: A27699; MUID:88209021; PMID:3365237<br>A; Accession: A27699                                                                                                                                                                                                                                                                                                                                                                                                           |

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A Accession: A01153; MUDI:86030270; PMID:4054110

A AAccession: A01153

A AAccession: A01154

A ACCESSION: A01154

A ACCESION: A01154

A ACCESSION: A01154

A ACCESSION: A01154

A ACCESSIO
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A; Residues: 'ANSM', 253-377 <AUE>
A; Note: dliferences are due to known cloning artifacts
B; Lottspelch, F;; Kellermann, J; Henschen, A.; Foertsch, B.; Muller-Esterl, W.
Eur. J. Biochem. 152, 307-314, 1985
A; Title: The amino acid sequence of the light chain of human high-molecular-mass kini
A; Reference number: A91153; MUID:86030270; PMID:4054110
   A. Accession: A34030
A. Molecule type: protein
A. Residues: 380-389 <-Sas.
A. Residues: 380-389 <-Sas.
A. Tenarcic, B.; Gabrijelcic, D.; Rozman, B.; Drobnic-Kosorok, M.; Turk, V.
Biol. Chem. Hoppe-Seyler 369, 257-261, 1988
A. Title: Human cathepsin B and cysteine proteinase inhibitors (CPIs) in inflammatory
A. Reference number: S02482; MUID:89076517; PMID:3264507
A. Molecule type: protein
A. Molecule type: protein
A. Molecule type: protein
A. Residues: 1-19;189-192; 310-314;381-389 <-LENI>
BEBS Lett. 232, 252-254, 1988
A. Title: Isolation and identification of hydroxyproline analogues of bradykinin in hu
A. Reference number: A61495; MUID:88211869; PMID:3366244
      FEBS Lett. 321, 93-97, 1993
A;Title: Cloning, expression and characterization of human kininogen domain
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A; Molecule type: protein
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A; Note: this peptide had Pro-383 modified to 4-hydroxyproline
A; Note: this peptide had Pro-383 modified to 4-hydroxyproline
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F;142-253/Domain: cystatin homology <CY2>
F;181-375/Domain: cystatin homology <CY2>
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F;383/Modified site: A-hydroxyproline (Pro) (partial) #status experimental
F;389-390/Cleavage site: Arg-Ser (kallikrein) #status experimental
F;401/Binding site: carbohydrate (Thr) (covalent) #status absent
  Nichinogen, HWW precursor [validated] - human
Nichinogen, HWW precursor [validated] - human
Nichinogen, HWW precursor [validated] - human
Nichinogen III hum kininogen III low molecular We
Cispecies: Homo sapiens (man)
Cispecies: Homo sapiens
Cisp
  A;Contents: annotation; gene organization
B;Plerce, J V.
B;Plerce,
  gene and a model for its evoluti
             s,
R;Kitamura, N.; Kitagawa, H.; Fukushima, D.; Takagaki, Y.; Miyata, T.; Nakanishi, J. Biol. Chem. 250, 8610-8617, 1985
A;Title: Structural organization of the human kininogen gene and a model for its e A;Reference number: A92545; MUID:85234583; PMID:2989294
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C;Accession: A25486
R;Kitagawa, H.; Kitamura, N.; Hayashida, H.; Miyata, T.; Nakanishi, S.
D; Biol. Chem. 262, 2190-2198, 1987
A;Title: Differing expression patterns and evolution of the rat kininogen gene family A;Reference number: A92625; MUID:87137443; PMID:3029068
  R.Furuto-Kato, S.; Matsumoto, A.; Kitamura, N.; Nakanishi, S.
J. Biol. Chem. 260, 12054-12059, 1985
A;Title: Primary structures of the mRNAs encoding the rat precursors for bradykinin a
   N.Contains: bradykinin
C.Species: Rattus norvegicus (Norway rat)
C.Date: 08-Mar-1989 #sequence_revision 08-Mar-1989 #text_change 15-Nov-1996
   C;Species: Rattus norvegicus (Norway rat)
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C;Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 18-Feb-2000
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R;Wilkinson, J.
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   1 NNATFYFK 8
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  A; Accession: A28055
  kininogen, HMW I
   Query Match
  Query Match
   Matches
   RESULT 5
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F;142-253/Domain: cystatin numblogy <CY2>
F;264-375/Domain: cystatin homology <CY2>
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A; Molecule type: protein

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R; Rittemura, N: Xitaqawa, H.; Fukushima, D.; Takagaki, Y.; Miyata, T.; Nakanishi, S.

J. Biol. Chem. 260, 8610-8617, 1985

J. Biol. Chem. 260, 8610-8617, 1985

J. Hitle: Structural organization of the human kininogen gene and a model for its evoluti

A; Reference number: A92545; MUID:85234583; PMID:2989294

A; Contents: annotation; gene organization
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A; Molecule type: protein
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Biochem. J. 307, 341-346, 1995
Biochem. J. 307, 341-346, 1995
A; Title: Human mast cell tryptase isoforms: separation and examination of substrate-spec A; Reference number: 555239; MUID:95251593; PMID:7733867
A; Accession: S55239
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A; Residues: 450-452, 7x, Maachl, F.; le Nguyen, D.; Becchi, M.; Heulin, M.H.; Nabet, P.; Bellevil
FEBS Lett. 373, 207-211, 1995
A; Title: Purification from human plasma of a tetrapeptide that potentiates insulin-like
A; Reference number: 568059; MUID: 96033974; PMID: 7589467
A; Accession: 568059
  ö
R;Lenarcic, B.; Krasovec, M.; Ritonja, A.; Olafsson, I.; Turk, V.
FEBS Lett. 280, 211-215, 1991
A;Title: Inactivation of human cystatin C and kininogen by human cathepsin D.
A;Reference number: S14303; MUID:91192133; PMID:2013314
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  KiPlerce, J.Y.
Fed. Proc. 27, 52-57, 1968
Affitle: Structural features of plasma kinins and kininogens.
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submitted to the EMBL Data Library, November 1995

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C; Accession: B83803
R; Takami, H; Nakasone, K; Takaki, Y; Maeno, G; Sasaki, R.; Masui, N.; Fuji, F.; H
Nucleic Acids Res. 28, 4317-4000
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R; Davies, R.M.; Devlin, K.; Duthoy, S.; Feltwell, T.; Fraser, A.; Hamlin, N.; Holro eam, M.A.; Rutherford, K.M.
Nature 409, 1007-1011, 2001
A;Authors: Rutter, S.; Seeger, K.; Simon, S.; Simmonds, M.; Skelton, J.; Squares, R.; A;Title: Massive gene decay in the leprosy bacillus.
A;Reference number: A86909; MUID:21128732; PMID:11234002
   ö
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  ABA-induced plasma membrane protein PM 19 - wheat
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Submitted to the EMBL Data Library, November 1996
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   2 NATFYF 7
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   C: Genetics
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R;Bryan, T.M.; Sperger, J.M.; Chapman, K.B.; Cech, T.R.
Proc. Natl. Acad. Sci. U.S.A. 95, 8479-8484, 1998
A;Title: Telomerase reverse transcriptuse genes in Tetrahymena thermophila and Oxytriche A;Reference number: 220985; MUID:98337940; PMID:9671703
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  Length 182;
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R; Theologis, A.; Ecker, J.R.; Palm, C.J.; Federspiel, N.A.; Kaul, S.; White, O.; Alon Chin, C.W.; Chung, M.K.; Conn, L.; Conway, A.B.; Conway, A.R.; Creasy, T.H.; Dewar, ansen, N.F.; Hudjes, B.; Huizar, L.
Nature 408, 816-820, 2000
A; Authors: Hunter, J.L.; Jenkins, J.; Johnson-Hopson, C.; Khan, S.; Khaykin, E.; Kim, A; Authors: Hunter, J.L.; Jenkins, J.; Johnson-Hopson, C.; Khan, S.; Khaykin, E.; Kim, Rizzo, M.; Rooney, T.; Rowley, X.; Liu, S.X.; Liu, Z.A.; Luros, J.S.; Maiti, R.; Maizia Rizzo, M.; Rooney, T.; Rowley, D.; Sakano, H.
A; Authors: Salzberg, S.L.; Schwartz, J.R.; Shinn, P.; Southwick, A.M.; Sun, H.; Tallo ker, M.; Wu, D.; Yu, G.; Fraser, C.M.; Venter, J.C.; Davis, R.W.
A; Title Sequence and analysis of chromosome 1 of the plant Arabidopsis.
A; Reference number: A86141; MUID:21016719; PMID:11130712
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C;Genetics:
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   probable cytochrome P450 T8L23.21 [imported] - Arabidopsis thaliana C;Species: Arabidopsis thaliana (mouse-ear cress)
C;Date: 02-Mar-2001 #sequence_revision 02-Mar-2001 #text_change 23-Mar-2001 C;Accession: G96611
   hypothetical protein MO1A10.4 - Caenorhabditis elegans
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C;Accession: T25830
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   R)Scheet, P. submitted to the EMBL Data Library, February 1997 A;Description: The sequence of C. elegans cosmid M01Al0. A;Reference number: Z20094
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Pred. No. 68;
1; Mismatches
   Score 34; DB 2;
Pred. No. 54;
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Best Local Similarity 75.0%;
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   1 NNATFYFK 8
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  A; Accession: T25830
   RESULT 15
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  Figolotin, A. Wincker, P.; Mauger, S.; Jaillon, O.; Malarme, K.; Weissenbach, J.; Ehrli Genome Res. 11, 731-753, 2001
A;Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis se A;Reference number: A86625; MUID:21235186; PMID:11337471
A;Accession: A86650
A;Status: preliminary
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C;Genetics:
   NADH2 dehydrogenase (ubiquinone) (EC 1.6.5.3) chain 2 - Pedinomonas minor mitochondrion C.Species: mitochondrion Pedinomonas minor C.Species: mitochondrion Pedinomonas minor C.Species: mitochondrion Pedinomonas minor C.Species: mitochondrion Pedinomonas minor C.Species: mitochondrion 16-Jul-1999 #text_change 03-Jun-2002 C.Species: mitochondrion 16-Jul-1999 #text_change 03-Jun-2002 C.Species: mitochondrion 16-Jul-1999 #text_change 03-Jun-2002 C.Species: M.W. R.Turmel, M.; Lemieux, C.; Burger, G.; Lang, B.F.; Otis, C.; Plante, I.; Gray, M.W. Submitted to the EMBL Data Library, December 1998 A.Spescription: The complete mitochondrial DNA sequences of Nephroselmis olivacea and Ped A.Reference number: 217261
  rhamnosyltransferase [imported] - Lactococcus lactis subsp. lactis (strain IL1403)
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  1 NNATEYE 7
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A;Gene: nad2
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R; Henkhaus, J; Wohldmann, P; Beck, C.
Submitted to the EMBL Data Library, July 1997
A; Description: The sequence of C. elegans cosmid C49D10.
A; Reference number: 221108
A; Reference number: 221108
A; Reference number: 221108
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A; Experimental source: strain Bristol N2; clone C49D10
C; Genetics:
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598 SNAQFYFK 605
   1 NNATFYFK 8
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Search completed: March 13, 2003, 19:14:01 Job time : 12 secs

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  saccharomyc
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MEDLINE-85122621; PubMed=6441591;
Ohkubo I., Kurachi K., Takasawa T., Shiokawa H., Sasaki M.;
Ohkubo I., Rurachi K., Takasawa T., Shiokawa H., Sasaki M.;
Isolation of a human cDNA for alpha 2-thiol proteinase inhibitor and
its identity with low molecular weight kininogen.";
Biochemistry 23:5691-5697(1984).
   Takagaki Y., Kitamura N., Nakanishi S.; "Cloning and sequence analysis of cDNAs for human high molecular weight and low molecular weight prekininogens. Primary structures of two human prekininogens."; J. Biol. Chem. 260:8601-8609(1985).
   Nakanishi S.; "Structural organization of the human kininogen gene and a model for
  amino acid sequence of the light chain of human high-molecular-kininogen.";
  P01042; P01043;
21-7UL-1986 (Rel. 01, Created)
01-FEB-1996 (Rel. 33, Last sequence update)
16-0CT-2001 (Rel. 40, Last annotation update)
Kininogen precursor (Alpha-2-thiol proteinase inhibitor) (Contains:
   Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
  Kitamura N., Kitagawa H., Fukushima D., Takagaki Y., Miyata T.,
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P34845
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   MEDLINE-86030270; PubMed-4054110;
Lottspeich F., Kellermann J., Henschen A., Foertsch B.,
Mueller-Esterl W.;
  Pierce J.V.; "Structural features of plasma kinins and kininogens."; Fed. Proc. 27:52-57(1968).
  Sueyoshi T., Miyata T., Kato H., Iwanaga S.; "Disulfide bonds in bovine HWW kininogens.";
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CPBJ_MOUSE
YIV9_YEAST
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  CHLE_HUMAN
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   Eur. J. Biochem. 152:307-314(1985)
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  GENE STRUCTURE.
   its evolution.
  Bradykinin].
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   saccharomyc
  saccharomyc
pseudalatia
   oxytricha t
  mus musculu
  heliothis a
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  Lostridium
  buchnera ap
  homo sapien
   rattus norv
   sapien
   escherichia
   vibrio chol
   bos taurus
h platelet-
   trichoplusi
   dictyosteli
   escherichia
   methanococc
   acinetobact
   helicobacte
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   bos taurus
  salmonella
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  drosophila
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  OMPU_VIBCH
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VEF_GVPU
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VEF_GVHA
ST23_YEAST
NUC2_NEUCR
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NIXA_HELPJ
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Maximum Match 100%
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  Minimum DB
Maximum DB
   Database :
   Sequence:
  Searched:
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RESULT 2
   KNG_RAT
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   This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation her European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
  -i- SUBCELLULAR LOCATION: Secreted.
-i- ALTERNATIVE PRODUCTS: 2 ISOFORMS; HMW (SHOWN HERE) AND LAWW; ARE PRODUCED BY ALTERNATIVE SPLICING.
-i- TISSUE SPECIFICITY: PLASMA.
-i- PTM: BRADYKININ IS RELEASED FROM KININOGEN BY PLASMA KALLIKREIN.
-i- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
  Glycoprotein; Plasma; Repeat; Thiol protease inhibitor; Vasodilator; Bradykinin; Blood coagulation; Inflammatory response; Signal;
  KININOGEN
   JOINED.
JOINED.
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JOINED.
  JOINED.
   JOINED.
   JOINED.
  InterPro; IPR000010; Cystatin.
InterPro; IPR002395; Kininogen.
Pfam; PF00031; cystatin; 3.
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   Rattus norvegicus (Rat).
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Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
NCBI_TaxID=10116;
   Gaps
            BRADYKININ.
KININGEN LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
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ACTIVITY).
  Kitagawa H., Kitamura N., Hayashida H., Miyata T., Nakanishi S.; "Differing expression patterns and evolution of the rat kininogen gene family.";
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Furuto-Kato S., Matsumoto A., Kitamura N., Nakanishi S.;
  -> I (IN REF. 1).
3132B4CBAF8FBB7E CRC64;
KININGEN HEAVY CHAIN
   (GLCNAC.
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  KNG_RAT STANDARD; PRT; 639 AA. P08934; P08933; 01-NOV-1988 (Rel. 09, Created) 01-NOV-1988 (Rel. 09, Last sequence update) 16-OCT-2001 (Rel. 40, Last annotation update) Kininogen precursor [Contains: Bradykinin].
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  DOMAIN
  DOMAIN
   REPEAT
REPEAT
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                            CHAIN
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   TISSUE SPECIFICITY: PLASMA.

THE SADDYKININ IS RELEASED FROM KININGEN BY PLASMA KALLIKREIN.

MISCELLANEOUS: RAT EXPRESS FOUR TYPES OF KININGENS: THE CLASSICAL

HWW/LMW KININGENS AND TWO ADDITIONAL LMW-LIKE KININGENS: TI AND
   "Structure and expression of the genes for major acute phase alpha 1-protein (thiostatin) and kininogen in the rat.";
J. Biol. Chem. 262:9298-9308(1987).
  FIRST AZ8023, AZ8025.
InterPro; IPR000010; Cystatin.
InterPro; IPR0003195; Kininogen.
Pfam; PF00031; cystatin; 3.
PRINTS; PR00334; KININOGEN.
SMART; SM00043; CY; 3.
PROSITE; PS00287; CYSTATIN; 2.
Glycoprotein; Plasma; Repeat; Thiol protease inhibitor; Vasodilator; Bradykinin; Blood coagulation; Inflammatory response; Signal; SIGNAL

1 18
   SUBCELLULAR LOCATION: Secreted.
ALTERNATIVE PRODUCTS: 2 ISOFORMS; HWW (SHOWN HERE) AND LAW; ARE PRODUCED BY ALTERNATIVE SPLICING.
"Primary structures of the mRNAs encoding the rat precursors for bradykinin and T-kinin. Structural relationship of kininogens with major acute phase protein and alpha 1-cysteine proteinase
   SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
  Biol. Chem. 260:12054-12059(1985).
  EMBL; M14369; AAA41484.1; -.
EMBL; M14369; AAA41485.1; ALT_SEQ.
EMBL; M16455; AAA41482.1; -.
PIR; A25466; A25486.
PIR; A28055; A28055.
  STRAIN-Wistar; TISSUE-Liver;
MEDLINE-87137465; PubMed-3818598;
  STRAIN-Buffalo;
MEDLINE-87250580; PubMed-2439509;
Fung W.-P., Schreiber G.;
  EMBL; L29428; AAA41486.1; -. EMBL; M11884; AAA41487.1; -.
   SEQUENCE OF 1-65 FROM N.A.
  SEQUENCE OF 1-41 FROM N.A.
   CLOTTING
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   N-LINKED (GLCNAC. .) (POTENTIAL).
N-LINKED (GLCNAC. .) (POTENTIAL).
N-LINKED (GLCNAC. .) (POTENTIAL).
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N-LINKED (GLCNAC. .) (POTENTIAL).
N-LINKED (GLCNAC. .) (POTENTIAL).
N-LINKED (GLCNAC. .) (POTENTIAL).
VEPSYIARVÖERRDEGNEGOFIHGHGWILHAKO -> RLLNS
CEYKGRLLKAGAGPAPERQAEASTVTP (IN ISOPORM
   Bryan T.M., Sperger J.M., Chapman K.B., Cech T.R.;
"Telomerase reverse transcriptase genes identified in Tetrahymena thermophila and Oxytricha trifallax.";
Proc. Natl. Acad. Sci. U.S.A. 95:8479-8484(1998).
-I- FUNCTION: TELOMERASE IS A RIBONUCLEOPROTEIN ENZYME ESSENTIAL FOR THE REPLICATION OF CHROMOSOME TERMINI IN MOST BUKARYOTES. IT ELONGATES TELOMERES. IT IS A REVERSE TRANSCRIPTASE THAT ADDS SIMPLE SEQUENCE REPEARTS TO CHROMOSOME ENDS BY COPYING A TEMPLATE SEQUENCE MITHIN THE RNA COMPONENT OF THE ENZYME.
   Gaps
   TERT_OXYTR STANDARD; PRT; 1132 AA.
076332;
30-MAY-2000 (Rel. 39, Created)
30-MAY-2000 (Rel. 39, Last sequence update)
30-MAY-2000 (Rel. 39, Last annotation update)
Telomerase reverse transcriptase (EC 2.7.7.-) (Telomerase catalytic
   ö
   Eukaryota; Alveolata; Ciliophora; Spirotrichea; Stichotrichia;
Stichotrichida; Oxytrichidae; Oxytricha
   SUBCELLULAR LOCATION: Nuclear.
SIMILARITY: BELONGS TO THE REVERSE TRANSCRIPTASE FAMILY.
  86.7%; Score 39; DB 1; Length 639; 87.5%; Pred. No. 3.7; tive 0; Mismatches 1; Indels
   INTERCHAIN (BY SIMILARITY).
BY SIMILARITY.
  LMW).
MISSING (IN ISOFORM LMW).
  -> K (IN REF. 2).
D3172DF94FF56AF5 CRC64;
                                   KININGEN LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 3.
HIS-RICH.
             KININOGEN HEAVY CHAIN.
   subunit) (Telomerase subunit P133)
  SEQUENCE FROM N.A. MEDLINE-98337940; Pubmed-9671703;
   434 639 M
61 61 E
639 AA; 70933 MW;
   TELOMERASE SUBFAMILY.
   7; Conservative
  Oxytricha trifallax.
  Query Match
Best Local Similarity
  NCBI_TaxID=5946;
   1 NNATFYFK 8
CONFLICT
   DISULFID
  CARBOHYD
   DISULFID
  DISULFID
  DISULFID
   DISULFID
   DISULFID
  DISULFID
  DISULFID
   CARBOHYD
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   CARBOHYD
   CARBOHYD
   CARBOHYD
   VARSPLIC
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  DOMAIN
DOMAIN
   DOMAIN
   DOMAIN
                                       CHAIN
   TERT_OXYTR
   Matches
  RESULT 3
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InterPro; IPR000997; Cholinesterase
   SIGNAL
  CHAIN
  INVG_SALTY
   Matches
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   and for commercial
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  entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
   Rachinsky T.L., Camp S., Li Y., Ekstroem T.J., Newton M., Taylor P.; "Molecular cloning of mouse acetylcholinesterase: tissue distribution of alternatively spliced mRNA species."; Newton 5:317-327(1990).
   SIMILARITY: BELONGS TO THE TYPE-B CARBOXYLESTERASE/LIPASE FAMILY.
  Gaps
   EMBL; AF060230; AAC39163.1; -.
InterPro; IPR003545; Telomerase_RT.
PRINTS; PR01365; TELOMERASERT.
Transferase; RNA-directed DNA polymerase; Telomere; Nuclear protein;
   01-OCT-1993 (Rel. 27, Created)
01-OCT-1993 (Rel. 27, Last sequence update)
15-UUL-1998 (Rel. 36, Last annotation update)
Cholinesterase precursor (EC 3.1.1.8) (Acylcholine acylhydrolase)
(Choline esterase II) (Butyrylcholine esterase)
   Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus
   ;
  Score 36; DB 1; Length 1132;
Pred. No. 24;
1; Mismatches 1; Indels
  DNA-binding.
SEQUENCE 1132 AA; 134124 MW; 01E145F5F24392DC CRC64;
   Usage by
modified and this statement is not removed. entities requires a license agreement (See hor send an email to license@isb-sib.ch).
   603 AA
   PIR, A39768, A39768.
HSSP, P21836, 1MAH.
MGD, MG18942PB, Bche
InterPro; IPR002018; CarbesteraseB.
  SEQUENCE FROM N.A. MEDLINE-90380429; PubMed-2400605;
  80.0%;
  SEQUENCE OF 97-237 FROM N.A.
   EMBL; M99492; AAA37328.1; -
   6; Conservative
   STANDARD;
  Pseudocholinesterase)
  Mus musculus (Mouse).
  1001 NNISFYFK 1008
   Query Match
Best Local Similarity
   NCBI_TaxID-10090;
  œ
  1 NNATFYFK
  TISSUE-Liver;
  CHLE_MOUSE
  003311;
  CHLE_MOUSE
  Matches
   RESULT
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MEDLINE=21534948; PUDMed=11677609; McClelland M., Sanderson K.E., Spieth J., Clifton S.W., Latreille P., Courtney L., Porwollik S., Ali J., Dante M., Du F., Hou S., Layman D., Leonard S., Nguyen C., Scott K., Holmes A., Grewal N., Mulvaney E., Ryan E., Sun H., Florea L., Miller W., Stoneking T., Nhan M.,
  Gaps
  BY 5...
BY SIMILARITY.
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A 
  Salmonella typhimurium.
Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
   MEDLINE=95172719; PubMed=7868245;
Lodge J., Douce G.R., Amin I.I., Bolton A.J., Martin G.D.,
Chaffield S., Dougan G., Brown N.L., Stephen J.;
Blological and genetic characterization of InphoA mutants of
Salmonella typhimurium TML in the context of gastroenteritis.";
Infect. Immun. 63:762-769(1995).
   MEDLINE-95089692; PubMed-7997169;
Kaniga K., Bossio J.C., Galan J.E.;
"The Salmonella typhimurium invasion genes invF and invG encode
homologues of the AraC and PulD family of proteins.";
MOI. Microbiol. 13:555-568(1994).
  ö
   Score 33; DB 1; Length 603; Pred. No. 48;
   Indels
  719B1B220D1E5367 CRC64;
   Hydrolase; Serine esterase; Glycoprotein; Signal.
   (Rel. 41, Last sequence update)
(Rel. 41, Last annotation update)
  CHOLINESTERASE
   562 AA
   0; Mismatches
  PROSITE; PS00122; CARBOXYLESTERASE_B_1; 1. PROSITE; PS00941; CARBOXYLESTERASE_B_2; 1.
  SEQUENCE FROM N.A.
STRAIN-LT2 / SGSC1412 / ATCC 700720;
IPR000379; Ser_estrs_site.
  PRT;
   01-JUN-1994 (Rel. 29, Created)
15-JUN-2002 (Rel. 41, Last seq
                   Pfam; PF00135; COesterase; 1. PRINTS; PR00878; CHOLNESTRASE
   ..
X
   73.3%;
85.7%;
  68521
   6; Conservative
   STANDARD;
   484
510
  InvG protein precursor. INVG OR STM2898.
   603
2227
354
467
121
2292
548
  510
515
129
603 AA;
  Local Similarity
  SEQUENCE FROM N.A.
  SEQUENCE FROM N.A.
   ||| |||
443 NNAFFYF 449
  NCBI_TaxID=602;
  1 NNATFYF 7
  15-JUN-2002
  STRAIN-SR-11
  Salmonella.
  STRAIN-TML;
   INVG_SALTY
P35672;
InterPro;
  ACT_SITE
DISULFID
  DISULFID
DISULFID
   CONFLICT
   ACT_SITE
  ACT_SITE
  DISULFID
  CARBOHYD
   CARBOHYD
   CARBOHYD
   CARBOHYD
   CARBOHYD
  CARBOHYD
  CARBOHYD
   Query Match
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SEQUENCE FROM N.A.
STRAIN-S288c / AB972;
  GCN2 protein kinase
  Query Match
Best Local Similarity
Matches 5; Conserv
   SEQUENCE FROM N.A.
  SEQUENCE FROM N.A.
   505 NNTTYYLK 512
   NCBI_TaxID-4932;
  1 NNATFYFK 8
   GCN2_YEAST
  GCN2_YEAST
   RESULT 7
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  ö
   STRAIN-Landrace; TISSUE-Heart; MEDLINE-99365306; PubMed-10433971; Lin C.S., Sun Y.L., Liu C.Y., Yang P.C., Chang L.C., Cheng I.C., Mao S.J.T., Huang M.C.; "Complete nucleotide sequence of pig (Sus scrofa) mitochondrial genome and dating evolutionary divergence within articodactyla.";
                                 Nature 413:852-856(2001).
--- FUNCTION: INVOLVED IN THE INVASION OF THE CELLS OF THE INTESTINAL EPITHELIUM. COULD BE NECESSARY FOR THE EXPORT OF INVASION RELATED DETERMINANTS.
Waterston R., Wilson R.K.; ^{\prime} "Complete genome sequence of Salmonella enterica serovar Typhimurium
  Gaps
   RPVLLTQENVP -> APGITSSGKCS (IN REF. 2).
  Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
  EMBL; AE00883; AAL2178.1; -.
EMBL; AE00883; AAL2178.1; -.
StyGene; SG10188; invg.
InterPro; IPR004846; GSPII/IIIprotein.
InterPro; IPR004845; GSPII/IIIprotein.
InterPro; IPR00485; GSPII_III; 1.
PROSITE; PS00875; T2SP.D; 1.
Virulence; Transport; Protein transport; Signal; Outer membrane;
  A -> R (IN REF. 2).
E -> O (IN REF. 2).
LRDOKWIP -> CAIRWWER (IN REF. 2).
AMPAFSANG -> RCORFORM (IN REF. 2).
  ö
   -i- SUBCELLULAR LOCATION: Outer membrane (Potential).
   Score 32; DB 1; Length 562;
Pred. No. 70;
3; Mismatches 2; Indels
   8022905BE256058D CRC64;
  G -> S (IN REF. 2).

AAA -> KPAEQ (IN REF. 2)

S -> T (IN REF. 1 AND 2)

I -> V (IN REF. 2).
   15-JUN-2002 (Rel. 41, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
NADH-ubdquinone oxidoreductase chain 5 (EC 1.6.5.3).
MINDS OR NDS OR NADHS.
   606 AA.
   INVG PROTEIN.
   POTENTIAL
   ö
  NUSM_PIG STANDARD; E Q9TDR1; 15-JUN-2002 (Rel. 41, Created)
   61765 MW;
   EMBL; X75302; CAA53049.1; -. EMBL; U08280; AAA74040.1; -.
   71.1%;
ilarity 75.0%;
Conservative
   232
243
262
328
329
562 AA;
   Local Similarity
   SEQUENCE FROM N.A.
  Complete proteome
  385 NNRTEYTK 392
   Sus scrofa (Pig).
  NCBI_TaxID=9823;
  1 NNATEYEK 8
  Mitochondrion
   15-JUN-2002
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   CONFLICT
CONFLICT
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   CONFLICT
  Query Match
  CONFLICT
   SIGNAL
  Matches
                       LT2.
  RESULT 6
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  ö
   and histidyl-
   Roussou I., Thireos G., Hauge B.M.; "Transcriptional-translational regulatory circuit in Saccharomyces cerevisiae which involves the GCN4 transcriptional activator and the
  Gaps
  MEDLINE-89282814; PubMed-2660141; Wek R.C., Jackson B.M., Hinnebusch A.G.; Wackson B.M., Hinnebusch A.G.; Jackson B.M., Hinnebusch A.G.; Jackson Gomains homologous to protein kinases and histidy tRNA synthetases in GGN2 protein suggests a mechanism for coupling GCN4 expression to amino acid availability."; Proc. Natl. Acad. Sci. U.S.A. 86:4579-4583(1989).
   Saccharomyces cerevisiae (Baker's yeast).
Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
Saccharomycetales; Saccharomycetaceae; Saccharomyces.
Gene 236:107-114(1999).
-!- CATALYTIC ACTIVITY: NADH + ubiquinone = NAD(+) + ubiquinol.
  ö
   MEDLINE-91246169; PubMed-2038314;
Ramirez M., Wek R.C., Hinnebusch A.G.;
"Ribbosome association of GCN2 protein Kinase, a translational activator of the GCN4 gene of Saccharomyces cerevisiae.";
MOI. Cell. Biol. 11:3027-3036(1991).
  Score 32; DB 1; Length 606; Pred. No. 75;
  Indels
  Submitted (MAR-1996) to the EMBL/GenBank/DDBJ databases
  Pfam; PF00361; oxidored_q1; 1.
Pfam; PF00662; oxidored_q1_N; 1.
Oxidoreductase; NAD; Ubiquinone; Mitochondrion.
SEQUENCE 606 AA; 68649 MW; F5D6C256A241D284 CRC64;
  .;
;
  01-NOV-1991 (Rel. 20, Last sequence update) 30-MAY-2000 (Rel. 39, Last annotation update)
   PRT; 1590 AA
  Pred. No. 75;
1; Mismatches
   InterPro; IPR001750; Oxidored_q1.
InterPro; IPR001516; Oxidored_q1_N.
   Mol. Cell. Biol. 8:2132-2139(1988)
  Protein kinase GCN2 (EC 2.7.1.-). GCN2 OR AAS1 OR YDR283C.
   MEDLINE-88261291; PubMed-3290651;
   ACTIVITY ON SUI2.
MEDLINE=92154672; PubMed=1739968;
  01-APR-1990 (Rel. 14, Created)
  71.18;
62.58;
   EMBL; AF034253; AAD34195.1;
  ASSOCIATION WITH RIBOSOMES
  5; Conservative
   STANDARD;
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  Buchnera aphidicola (subsp. Acyrthosiphon pisum) (Acyrthosiphon pisum
   16-6CT-2001 (Rel. 40, Created)
16-6CT-2001 (Rel. 40, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
1midazole glycerol phosphate synthase subunit hisH (EC 2.4.2.-) (IGP synthase glutamine amidotransferase subunit) (IGP synthase subunit
  Sus scrofa (Pig).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Suina; Suidae; Sus.
NCBI_TaxID=9823;
  Nuclear protein; RNA-binding; mRNA splicing; Zinc-finger; Repeat.
DOMAIN 47 >82 INVOLVED IN BINDING UZAF-65 (BY
01-NOV-1997 (Rel. 35, Created)
01-NOV-1997 (Rel. 35, Last sequence update)
15-UUN-2002 (Rel. 41, Last annotation update)
Splicing factor U2AF 35 kDa subunit (U2 auxiliary factor 35 kDa subunit) (U2 snRNP auxiliary factor small subunit) (Fragment).
U2AF1.
  Score 31; DB 1; Length 82;
Pred. No. 15;
1; Mismatches 1; Indels
  hisH) (ImGP synthase subunit hisH) (IGPS subunit hisH).
HISH OR BU103.
   Bacteria; Proteobacteria; gamma subdivision; Buchnera
  96882458044823A8 CRC64;
  SIMILARITY).
   InterPro; IPR000571; Znf_CCCH. Pfam; PF00642; zf-CCCH; 1.
  68.9%;
71.4%;
   EMBL; F14757; CAA23231.1; -.
  9416 MW;
   Conservative
   symbiotic bacterium)
   Similarity 5; Conserv
   SEQUENCE FROM N.A.
  82 AA;
   SEQUENCE FROM N.A.
   NCBI_TaxID-118099;
   17 NCSFYFK 23
   2 NATFYFK 8
  SEQUENCE
  HIS5_BUCAI
  Query Match
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  HIS5_BUCAI
   Matches
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   ő
   -i- FUNCTION: STIMULATES GCN4 TRANSLATION IN AMINO ACID-STARVED CELLS BY PHOSPHORYLATING THE ALPHA SUBUNIT OF EIF-2 (SUI2) ON SER-52.
-i- SUBCELLULAR LOCATION: ASSOCIATES WITH THE 60S SUBUNIT OF
                              "Phosphorylation of initiation factor 2 alpha by protein kinase GCN2 mediates gene-specific translational control of GCN4 in yeast."; Cell 68:585-596(1992).
  CAUTION: REF. 2 SEQUENCE DIFFERS FROM THAT SHOWN IN POSITIONS 301
   Gaps
   RIBOSOMES.
-1- INDUCTION: BY AMINO ACID STARVATION AND IN THE PRESENCE OF THE
  PROTEIN KINASE 1.
HISTIDYL-TRNA SYNTHETASE LIKE.
ATP (BY SIMILARITY).
ATP (BY SIMILARITY).
BY SIMILARITY).
TIKAKLP -> NYKOKIA (IN REF. 2).
LASSEMMEN -> VYFSNHGKS (IN REF. 2).
I -> M (IN REF. 2).
P -> A (IN REF. 2).
RERYASDEELAVL -> DHDMLLTLKRLHFS
  SIMILARITY: BELONGS TO THE SER/THR FAMILY OF PROFEIN KINASES. GCN2 SUBFAMILY.
   ö
  TO 364 AND FROM POSITION 981 ONWARD DUE TO FRAMESHIFTS.
   Score 32; DB 1; Length 1590;
Pred. No. 2e+02;
   PROSITE; PS50862; AA_TRNA_LIGASE_II; FALSE_NEG.
PROSITE; PS00107; PROTEIN_KINASE_ATP; 1.
PROSITE; PS50011; PROTEIN_KINASE_ST; 1.
PROSITE; PS50011; PROTEIN_KINASE_ST; 1.
Transferase; Serine/Chreonine-protein kinase; ATP-binding; Amino-acid biosynthesis; Aminoacyl_tRNA synthetase; Repeat.
  S -> T (IN REF. 2).

IL -> MI (IN REF. 2).

F -> C (IN REF. 2).

M -> K (IN REF. 2).

E -> Q (IN REF. 2).

My: 768815424F64FIAA CRC64;
Feng L., Wek R.C., Cigan A.M., Donahue T.D.,
   1; Indels
   82 AA.
  1; Mismatches
   (IN REF. 2)
S -> T (IN
  InterPro; IPR002106; AALRNA_ligasell.
InterPro; IPR000719; Euk_pkinase.
InterPro; IPR002290; Ser_thr_pkinase.
   EMBL; M27082; AAA34636.1; -.
EMBL; M20487; AAA34881.1; ALT_FRAME.
EMBL; U51030; AAB64461.1; ALT_INIT.
   PRT;
  ; Euk_pkinase; 3.
   71.1%;
75.0%;
   182041
   Conservative
   STANDARD;
  Pfam; PF00069; pkinase;
ProDom; PD000001; Euk_p
  658
770
885
   PIR; S05781; OKBYNZ.
PIR; A27723; A27723.
SGD; S0002691; GCNZ.
  553
571
658
770
885
1590 AA;
   Query Match
Best Local Similarity
Matches 6; Conserv
  11111 :1
266 NNATFVWK 273
   GCN4 PROTEIN
Dever T.E., Fen
Hinnebusch A.G.
  1 NNATFYFK 8
   RESULT 8
U2AG_PIG
ID U2AG_PIG
AC Q29350;
  NP_BIND
BINDING
ACT_SITE
CONFLICT
  CONFLICT
CONFLICT
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CONFLICT
SEQUENCE
   CONFLICT
CONFLICT
CONFLICT
CONFLICT
  CONFLICT
  DOMAIN
   DOMAIN
  <del>:</del>
   <u>:</u>
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Gaps

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196 AA.

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CONFLICT
   Query Match
  ZN_FING
DOMAIN
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   ö
  DECENTAIN—C57BL/63; TISSUE—Small intestine;

MEDLINE=2108560; PubMed=11217851;

KRWAI J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,

Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,

Alzawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,

Saito T., Okazaki Y., Gojobori T., Bono H., Kasukawa T., Saito R.,

Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,

Fleischmann W., Gasterland T., Gissi C., King B., Kochiwa H.,

Kuehl P., Lewis S., Matsuo Y., Nikaido I., Pesole G., Quackenbush J.,

Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
   Gaps
                MEDLINE-20445173; PubMed-10993077;
Shigenobu S., Watanabe H., Hattori M., Sakaki Y., Ishikawa H.;
"Genome sequence of the endocellular bacterial symbiont of aphids
   Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus
NCBI_TaxID=10090;
  InterPro; IPR000991; GATase_1.
Pfam: PF00117; GATase; 1.
PROSITE; PS00442; GATASE_TYPE_1; FALSE_NEG.
Histidine blosynthesis; Transferase; Glutamine amidotransferase;
   15-JUN-2002 (Rel. 41., Created)
15-JUN-2002 (Rel. 41, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
15-JUN-2002 (Rel. 41, Last annotation update)
5placing factor UZAF 35 kps subunit (UZ auxiliary factor 35 kps subunit) (UZ snRNP auxiliary factor small subunit).
   ö
  68.9%; Score 31; DB 1; Length 196; 71.4%; Pred. No. 37; 1.4%; Pred. 1; Mismatches 1; Indels
   SIMILARITY.
SIMILARITY.
904624B74BEBESD CRC64;
  BY SIMILARITY.
BY SIMILARITY.
BY SIMILARITY.
  239 AA
  PRT;
  77 77 BY
178 178 BY
180 180 BX
196 AA; 21679 MW;
  EMBL; AP001118; BAB12822.1; -.
   Query Match
Best Local Similarity 71.4%;
  STANDARD;
   Q9CZ98; Q99LX2;
  Mus musculus (Mouse).
   Complete proteome. ACT_SITE 77
  SEQUENCE FROM N.A.
  135 NNSRFYF 141
   1 NNATEYF 7
  RESULT 10
U2AG_MOUSE
ID U2AG_MOUSE
AC Q9D8B3; Q9CZ
   ACT_SITE
ACT_SITE
SEQUENCE
   UZAF1.
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  ID ACC OOC OOC OOC SERVER TO SERVER THE SERV
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  ö
Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G., Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F., Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M., Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H., Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P., Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N., Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F., Suzuki H., Toyo-oka K., Wang K.H., Weitz C., Whittaker C., Wilming L., Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
   Gaps
  Nuclear protein; RNA-binding; mRNA-splicing; Zinc-finger; Repeat. ZN_FING 13 41 C3H1-TYPE 1.
   Hayashizaki Y.; "Functional annotation of a full-length mouse cDNA collection.";
   ö
   Score 31; DB 1; Length 239;
Pred. No. 46;
   -> R (IN REF. 1; BAB25609)
DFF944210581244D CRC64;
  C3H1-TYPE 2. ARG/GLY/SER-RICH (RS DOMAIN)
  C3HL-TIPE 1.
RNA-BINDING (RRM).
   Mismatches
   EMBL; AK008332; BAB25609.1; --
EMBL; AK012849; BAB25513.1; --
EMBL; BC002184; AAH02184.1; --
MGD; MGI: 98884; UZafi...
InterPo: IPR000504; RRM_I.
InterPo: IPR00051; Znf_CCCH.
Pfam; PF00076; rrm; 2.
Pfam; PF00042; zf-CCCH; 4.
SMART; SM00361; RRM, 1.
  1;
   149 173 C3
179 238 AJ
187 187 G
239 AA; 27815 MW;
   68.98;
  71.48;
   SEQUENCE OF 2-239 FROM N.A.
   Nature 409:685-690(2001).
   5; Conservative
   TISSUE-Breast tumor;
  Best_Local Similarity
Matches 5; Conserv
   17 NCSFYFK 23
   NATFYFK 8
```

PIR; S27969; S27969. PIR; A46179; A46179. Genew; HGNC:12453; U2AF1. MIM; 191317; -.

```
InterPro; IPR000504; RNA_rec_mot.
InterPro; IPR000571; znf_CCCH.
Fram; PF00076; rrn; 1.
Pfam; PF00042; zf-CCH; 2.
SNART; SM00350; RRM; 1.
SNART; SM00356; znF_C311; 2.
  Ouery Match
Best Local Similarity 71.4،
انام 5; Conservative
        ð
  THE FUNCTION TO THE PERFORMANCE OF THE STATE   This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation-the Buropean Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
   MEDLINE=20289799; PubMed=10830953;

MEDLINE=20289799; PubMed=10830953;

MEDLINE=20289799; PubMed=10830953;

A Bark H.-S., Toyoda A., Ishilf K., Totoki Y., Choi D.-K., Soeda E.,

A Park H.-S., Toyoda A., Ishilf K., Totoki Y., Choi D.-K., Soeda E.,

A Nemacal U., Delabar J., Kumpf K., Lehmann R., Patterson D.,

Rosenthal A., Rump A., Schillhabel M., Schudy A., Zimmermann W.,

Rosenthal A., Sasaki T., Nagamine K., Mitsuyama S., Antonarakis S.E.,

A Shiftani A., Sasaki T., Nagamine K., Mitsuyama S., Antonarakis S.E.,

Minoshima S., Shimizu N., Nordsiek G., Hornischer K., Brandt P.,

A Scharfe M., Schoen O., Desario A., Reichelt J., Kauer G., Bloccker H.,

Ramser J., Beck A., Klages S., Hennig S., Riesselmann L., Dagand E.,

Nemmseyer S., Borzym K., Gardiner K., Nizetic D., Francis F.,

Lehrach H., Reinhardt R., Yaspo M.-L.;

"The DNA sequence of human chromosome 21.";
  Zhang M., Zamore P.D., Carmo-Fonseca M., Lamond A.I., Green M.R., Cloning and intracellular localization of the U2 small nuclear ribonucleoprotein auxiliary factor small subunit.";
Proc. Natl. Acad. Sci. U.S.A. 89:8769-8773(1992).
   Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
   01-APR-1993 (Rel. 25, Created)
01-APR-1993 (Rel. 25, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
5-JUN-2002 (Rel. 41, Last annotation update)
Splicing factor "The subunit (U2 auxillary factor 35 kDa
   SEQUENCE FROM N.A., AND SEQUENCE OF 68-89 AND 126-151
   subunit) (U2 snRNP auxiliary factor small subunit). U2AF1 OR U2AF35.
   240 AA.
  PRT;
   TISSUE-Fetal brain;
MEDLINE-92409598; PubMed=1388271;
  Nature 405:311-319(2000).
  STANDARD;
  Homo sapiens (Human).
   NCBI_TaxID=9606;
   U2AG_HUMAN
  FUNCTION
                        J2AG_HUMAN
RESULT 11
```

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Gaps

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1; Length 240; 1; Indels

Score 31; DB Pred. No. 46; Pred. No. 46; 1; Mismatches

2 NATFYFK 8

PROSITE; PS50102; RRM; 1. PROSITE; PS00030; RRM, RNP\_1; FALSE NEG. Nuclear protein; RNA-binding; mRNA splicing; Zinc-finger; Repeat. ZN\_FFNG 13 41

C3H1-TYPE 1. RNA-BINDING (RRM).

C3H1-TYPE 2

ARG/GLY/SER-RICH (RS DOMAIN). POLY-GLY.

3DA130DCE0B953F6 CRC64;

27872 MW; 68.98; 71.48;

SEQUENCE

ZN\_FING DOMAIN DOMAIN DOMAIN

```
Adams M.D., Celniker S.E., Holf R.A, Favans C.A., Gocayne J.D.,
RA Adams M.D., Celniker S.E., Holf R.A,
Racing R.A., Lewis S.E., Holf R.A,
Racing R.A., Lewis S.E., Stocker S., Ashburner M., Henderson S.N.,
RA Sutton G.G., Wortman J.R., Yandell M.D., Zhang Q., Chen L.X.,
RA Sutton G.G., Racter E.G., Helt G., Champe M., Pfelifer B.D.,
RA Man K.H., Doyle C., Baxter E.G., Helt G., Nelson C.R., Miklos G.L.G.,
RA Abril J.F., Agbayani A., An H.-J., Andrews-Pfannkoch C., Baldwin D.,
RA Ballew R.M., Basu A., Baxendale J., Bayraktaroglu L., Beasley E.M.,
Beeson K.Y., Benos P.V., Berman B.P., Bhandari D., Bolshakov S.,
RA Borkova D., Botchan M.R., Bouck J., Brokstein P., Brottier P.,
RA Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
RA Cherry J.M., Cawley S., Dahlke C., Davenport L.B., Davies P.,
RA Dodson K., Doup L.E., Downes M., Dugan-Rocha S., Dunkor B.C., Dunn P.,
RA Dorbin K.J., Evangelista C.C., Ferraz C., Ferriera S., Fleischmann W.,
RA Fosler C., Gabriellan A.E., Garg N.S., Gelbart W.M., Glasser K.,
RA Glodek A., Gong F., Gorrell J.H., Gu Z., Guan P., Harris M.,
   Drosophila melanogaster (Fruit fly).
Eukaryota; Metazoa; Arthropoda; Mandibulata; Pancrustacea; Hexapoda;
Insecta; Pterygota; Neoptera; Endopterygota; Diptera; Brachycera;
Muscomorpha; Ephydroidea; Drosophilidae; Drosophila.
   094535; 09VPN4;
01-NOV-1997 (Rel. 35, Created)
16-0CT-2001 (Rel. 40, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
59licing factor Uzaf 38 kDa subunit (UZ auxillary factor 38 kDa
subunit) (UZ snRNP auxillary factor small subunit).
UZAF38 OR CG3582.
  Rudner D.Z., Kanaar R., Breger K.S., Rio D.C.;
"Mutations in the small subunit of the Drosophila U2AF splicing factor cause lethality and developmental defects.";
Proc. Natl. Acad. Sci. U.S.A. 93:10333-10337(1996).
  264 AA.
  SEQUENCE FROM N.A., AND PARTIAL SEQUENCE.
  STRAIN-Berkeley;
MEDLINE-20196006; Pubmed-10731132;
   MEDLINE-96413646; PubMed-8816800;
  STANDARD;
   SEQUENCE FROM N.A.
   NCBI_TaxID=7227;
17 NCSFYFK 23
  U2AG_DROME
   RESULT 12
U2AG_DROME
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EMBL; M96982; AAA36619.1; -. EMBL; AP001748: BAA95534.1: -AP001748; BAA95534.1; us-09-461-061a-1.rsp

```
Harris N.L., Harvey D., Heiman T.J., Hernandez J.R., Houck J.,
Hostin D., Houston K.A., Howland T.J., Wei M.-H., Ibegwam C.,
Jaali M., Kalush F., Karpen G.H., Ke Z., Kennison J.A., Ketchum K.A.,
Kimmel B.E., Kodira C.D., Kraft C., Kravitz S., Kulp D., Lai Z.,
Lasko P., Lei Y., Levitsky A.A., Li J., Li Z., Liang Y., Lin X.,
Martei B.L., McIntosh T.C., McHeod M.P., McPherson D.,
Merkulov G., Milshina N.V., Mobarry C., Murris J., Moshrefi A.,
Mount S.M., Moy M., Murphy B., Murphy L., Murris J., Moshrefi A.,
Nelson D.K.,
Nelson K.A., Nixon K., Nusskern D.R., Parleson D.L.,
Palazzolo M., Pittman G.S., Pan S., Pollard J., Puri V., Reese M.G.,
Reinert K., Remington K., Saunders R.D.C., Scheeler F., Shen H.,
Spier E., Spradling A.C., Stapleton M., Strong R., Sunth T.,
Spier E., Spradling A.C., Stapleton M., Strong R., Sun E.,
Nung Z.-Y., Wassarman D.A., Weinstock G.M., Weissenbach J.,
Nang Z.-Y., Wassarman D.A., Weinstock G.M., Weissenbach J.,
Milliams S.M., Hoodage T., Worley K.C., Wu D., Yang S., Yao O.A.,
A. J., Yeh R.-F., Zaveri J.S., Zhan M., Zhang G., Zhao Q., Zheng L.,
Labeng X.H., Zhong F.N., Zhong W., Zhou X., Zhu X., Smith H.O.,
Science 287:218-2195(2000).
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   entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@lsb-sib.ch).
   ő
   I- FUNCTION: NECESSARY FOR THE SPLICING OF PRE-MRNA. BINDS TO THE POLYPYRIMIDINE TRACT OF INTRONS EARLY DURING SPLICEOSOME ASSEMBLY
  Gaps
   Zinc-finger; Repeat.
   ö
   SUBUNIT: ASSOCIATES WITH A 65 kDa PROTEIN (BY SIMILARITY).
  --- SUBCELLULAR LOCATION: NUCLEAR.
--- SIMILARITY: CONTAINS 1 RNA RECOGNITION MOTIF (RRM).
--- SIMILARITY: CONTAINS 2 C3H1-TYPE ZINC FINGERS.
--- SIMILARITY: BELONGS TO THE SR FAMILY OF SPLICING FACTORS.
   Score 31; DB 1; Length 264;
   Indels
  SMART; SMUGLOU, SMEN, 1.
PROSITE; PSS01023; RRM, 11.
PROSITE; PS00030; RRM_RNP 1; FALSE_NEG.
Nuclear protein; RNA-binding; mRNA-splicing; Zinc-finge
DOMAIN 44 149
RNA-BINDING (RRM).
180 213 ARG/SER-RICH (RS DOMAIN).
  -> D (IN REF. 1).
577285FB66FDB2F5 CRC64;
   334 AA.
   ARGC_ECOLI STANDARD; PRT; 334 AA P11446; 01-0CT-1989 (Rel. 12, Created) 01-0CT-1989 (Rel. 12, Last sequence update)
   50;
   Mismatches
   POLY-ARG.
POLY-GLY.
   Pred. No.
   EMBL; U67066; AAB17271.1; -.
EMBL; AE003590; AAF51512.1; -.
FLYBASSE; FR900017457; UZAF38.
INTERPRO; IPR000504; RRM_ISC_MOT.
INTERPRO; IPR000504; RRM_ISC_CCH.
Pfam; PF00076; ITM, 1.
Pfam; PF00642; Zf-CCH, 2.
SMART; SM00361; RRM_I; 1.
SMART; SM00356; ZRF_C3HI; 2.
  29877 MW;
   68.88;
  71.48;
   5; Conservative
   (BY SIMILARITY).
   264 AA;
   Local Similarity
   2 NATFYFK 8
   CONFLICT
  SEQUENCE
   Query Match
   DOMAIN
DOMAIN
DOMAIN
  RESULT 13
ARGC_ECOLI
  Matches
   Best
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   MEDLINE-83143275; PubMed-6761650;
Piette J., Cunin R., Boyen A., Charlier D.R.M., Crabeel M.,
van Vliet F., Glansdorff N., Squires C., Squires C.L.;
"The regulatory region of the divergent argECBH operon in Escherichia
  J. Bacteriol. 174:223-2331(1992).

-!- CATALYTIC ACTIVITY: N-acetyl-L-glutamate 5-semialdehyde + NADP(+)

- Phosphate = N-acetyl-5-glutamyl phosphate + NADPH.

- PATHMAY: Arginine biosynthesis; third step.

-!- PATHMAY: BELONGS TO THE NAGSA DEHYDROGENASE FAMILY.
   Meinnel T., Schmitt E., Mechulam Y., Blanquet S.; "Structural and blochemical characterization of the Escherichia coli
15-JUN-2002 (Rel. 41, Last annotation update)
N-acetyl-gamma-glutamyl-phosphate reductase (EC 1.2.1.38) (AGPR) (N.
acetyl-glutamate semialdehyde dehydrogenase) (NAGSA dehydrogenase).
  MEDLINE-89121510; PubMed-2851495; Barsot C., Boyen A., Cohen G.N., Glansdorff N.; Mayen A., Cohen G.N., Chan G.N., Cohen C.N.,   "Analysis of the Escherichia coli genome. IV. DNA sequence of the region from 89.2 to 92.8 minutes.";
   Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
Escherichia.
   Prodom; PD003765; AGPR_act_site; 1.
PROSITE; PS01224; ARGC; 1.
Arginine blosynthesis; Oxidoreductase; NADP; Complete proteome.
ACT_SITE 154
SEQUENCE 334 AA; 35952 MW; 67AC195ECEIC4789 CRC64;
  SEQUENCE FROM N.A.
STRAIN=K12 / MG1655;
MEDLINE=94089392; Pubmed=8265357;
Blattner F.R., Burland V.D., Plunkett G. III, Sofia H.J.,
  Nucleic Acids Res. 21:5408-5417(1993).
   Nucleic Acids Res. 10:8031-8048(1982).
   EMBL; AE000470; AAC76940.1; ...
PIR; JT0332; RDECEP.
ECGGENG; EG10065; argC.
Interpro; IPR000706; AGPR_act_site.
Interpro; IPR000534; Semialdh.
Pfam; PF01118; Semialdhyde_dh; 1.
Pfam; PF02774; Semialdhyde_dh; 1.
  EMBL; J01587; AABS9146.1; -.
EMBL; X55417; -; NOT_ANNOTATED_CDS.
EMBL; U00006; AAC43064.1; -.
   MEDLINE-92202162; PubMed-1551850;
   EMBL; M21446; AAA23477.1; -.
   SEQUENCE OF 1-19 FROM N.A.
  SEQUENCE OF 1-48 FROM N.A.
  SEQUENCE FROM N.A.
   argE gene product.
   Escherichia coli.
  NCBI_TaxID-562;
   OR B3958
   Daniels D.L.;
   STRAIN=K12
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CONFLICT
   KNL2_BOVIN
   Matches
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  SEQUENCE FROM N.A.

STRAIN-El Tor N16961 / Serotype Ol;

MEDLINE-20406833; PubMed=10952301;

Heidelberg J.F., Eisen J.A., Nelson W.C., Clayton R.A., Gwinn M.L.,

Dodson R.J., Haft D.H., Hickey E.K., Peterson J.D., Umayam L.A.,

Gill S.R., Nelson K.E., Read T.D., Tettelin H., Richardson D.,

Ermolaeva M.D., Vamathevan J., Bass S., Qin H., Dragoi I., Sellers P.,

McDonald L., Utterback T., Fleischmann R.D., Nierman W.C., White O.,

Salzberg S.L., Smith H.O., Colwell R.R., Mekalanos J.J., Venter J.C.,
                          Gaps
  Nature 406:477-483(2000).
  [1] SEQUENCE FROM N.A.
STRAIN-Classical Ogawa 395 / ATCC 39541 / Serotype 01;
STRAIN-STAIN-069; Pubmed-8945596;
Sperandio V., Bailey C.C., Giron J.A., Dirita V.J., Silveira W.D., Vettore A.L., Kaper J.B.;
  Bacteria; Proteobacteria; gamma subdivision; Vibrionaceae; Vibrio.
NCBI_TaxID=666;
  /ettore A.L., Kaper J.B.;
"Cloning and characterization of the gene encoding the OmpU outer
   'DNA sequence of both chromosomes of the cholera pathogen Vibrio
  InterPro; IPR003229; OMP_2.
InterPro; IPR001702; Porin_gram-ve.
Pfam; PF00267; Gram-ve_porins; 1.
Transmembrane; Porin; Signal; Outer membrane; Complete proteome.
                         ö
   STRAIN=El Tor;
ful Y., Zhang J.Z.;
"Cloning and expression of ompU of Vibrio cholerae and its
antigenicity analysis.";
  DB 1; Length 334;
  Submitted (APR-2000) to the EMBL/GenBank/DDBJ databases.
   OUTER MEMBRANE PROTEIN U.
  OMPU_VIBCH STANDARD; PRT; 341 AA. P97085; Q9L5A3; Q9KU90; 15-DEC-1998 (Rel. 37, Created) 16-OCT-2001 (Rel. 40, Last sequence update) 16-OCT-2001 (Rel. 40, Last annotation update) Outer membrane protein U precursor (Porin ompU)
  -> I (IN REF. -> K (IN REF.
           64;
                          Mismatches
 Score 31;
Pred. No.
  membrane protein of Vibrio cholerae.";
Infect. Immun. 64:5406-5409(1996).
  EMBL; AF253529; AAF64526.1; -.
EMBL; AE004149; AAF93799.1; ALT_INIT.
TIGR; VC0633; -.
68.9%;
75.0%;
  EMBL; U73751; AAB48973.1;
                         Conservative
  341
278
290
  [2]
SEQUENCE FROM N.A.
           Best Local Similarity
   109 NDATFYEK 116
   1 NNATFYFK 8
   Vibrio cholerae
  OMPU OR VC0633
  CHAIN
CONFLICT
CONFLICT
 Query Match
   cholerae
  SIGNAL
                       Matches
  RESULT 14
   셤
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   11 J. Biol. Chem. 262:2768-2779(1987).

21 J. Biol. Chem. 262:2768-2779(1987).

22 I. FUNCTION: (1) KININOGENS ARE INHIBITORS OF THIOL PROTEASES; (2)

23 LAW-KININOGEN INHIBITS THE AGERGATION OF THROMBOCYTES; (3) THE

24 VARIETY OF PHYSIOLOGICAL EFFECTS: (3A) INFLUENCE IN SMOOTH

25 A VARIETY OF PHYSIOLOGICAL EFFECTS: (3A) INFLUENCE IN SMOOTH

26 NATRIURESIS AND DIURESIS (KIDNEY).

27 I. SUBCELLOLLAR LOCATION: EXTRACELLULAR.

28 C. I. ALTERNATIVE PRODUCTS: 2 isoforms; LAW II (shown here) and HAW II

29 C. I. ALTERNATIVE PRODUCTS: 2 isoforms; LAW II (shown here) and HAW II

20 C. I. TISSUE SPECIFICITY: FLASMA.

21 ITSSUE SPECIFICITY: FLASMA.

22 C. I. PTM: BRADYKININ IS RELEASED FROM KININOGEN BY PLASMA KALLIKREIN.

23 C. I. PTM: BRADYKININ IS NELEASED FROM KININOGEN BY PLASMA KALLIKREIN.
  SHOMS
   Gaps
  21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
15-JUN-2002 (Rel. 41, Last annotation update)
Kininogen, LMW II precursor (Thiol proteinase inhibitor) (Contains:
Bradykinin].
   MEDLINE-83117859; PubMed=6572010;
Nawa H., Kitamura N., Hirose T., Asai M., Inayama S., Nakanishi S.;
"Primary structures of bovine liver low molecular weight kininogen precursors and their two mRNAs.";
Proc. Natl. Acad. Sci. U.S.A. 80:90-94(1983).
   MEDLINE-87137530; PubMed-3546295; Sueyoshi T., Miyata T., Hashimoto N., Kato H., Hayashida H., Miyata T., Iwanaga S.; Miyata T., Iwanaga S.; Bovine high molecular weight kininogen. The amino acid sequence, positions of carbohydrate chains and disulfide bridges in the heavy
  Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Bovoldea;
Bovidae; Bovinae; Bos.
  Glycoprotein; Plasma; Repeat; Vasodilator; Alternative splicing;
  ;
  DB 1; Length 341;
65;
  Indels
VG -> AS (IN REF. 1).
CECB39070E441732 CRC64;
  INVOLVED IN BLOOD CLOTTING.
-!- SIMILARITY: CONTAINS 3 CYSTATIN-LIKE DOMAINS.
   4.34 AA
  2; Mismatches
   Score 31;
Pred. No.
   EMBL; V00427; CAA23710.1; -. PIR; A01284; KGB012. HSP; P01038; 1A90.
InterPro; IPR000010; Cystatin. Pfam; PF00011; cystatin; 3. SMART; SM00043; CY; 3. PROSITE; PS00287; CXSTATIN; 2.
325 V
36645 MW;
   68.9%;
71.4%;
  5; Conservative
  STANDARD;
   Bos taurus (Bovine).
                               341 AA;
  Query Match
Best Local Similarity
   SEQUENCE FROM N.A.
  SEQUENCE OF 19-376.
  :||:|||
299 DATYYFK 305
   NCBI_TaxID=9913;
  chain portion."
  2 NATFYFK 8
  KNL2_BOVIN
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, 0,
   0; Gaps
   KININOGEN, LMW II.
HEANY CHAIN.
BRADKININ.
LIGHT CHAIN.
CYSTATIN-LIKE 1.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 2.
CYSTATIN-LIKE 6.
O-LINKED (GLCNAC. ..).
N-LINKED (GLCNAC. ..), OR 169.
N-LINKED (GLCNAC. ..); OR 169.
  Query Match 68.9%; Score 31; DB 1; Length 434; Best Local Similarity 62.5%; Pred. No. 83; Matches 5; Conservative 2; Mismatches 1; Indels
   48148 MW; 73A7079DE3E03430 CRC64;
Thiol protease inhibitor; Bradykinin; Signal.

SIGNAL

CHAIN

19
374
KININOGEN, LMW II.
CHAIN

BEPTIDE

CHAIN

ODMAIN

19
434
HEAVY CHAIN

BEPTIDE

CHAIN

387
434
HEAVY CHAIN

CHAIN

BOMAIN

19
525
CYSTATIN-LIKE 1.

CARBOHYD

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Search completed: March 13, 2003, 19:13:24 Job time : 6 secs

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Gaps
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"Complete genome sequence of the alkaliphilic bacterium Bacillus
"Complete genome sequence comparison with Bacillus subtilis.";
Nucleic Acids Res. 28:4317-4331(2000).
BEBL; AP001511; BAB04945.1;
Hypothetical protein; Complete proteome.
SEQUENCE 71 AA; 8026 MW; 311AC9AEB3C539D3 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
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STRAIN=RJ96BRP071;
Tanuti A., Swanson P.A., Devare S.G., Berro O.J., Savedra A.,
Tanuti A., Swanson P.A., Devare S.G., Berro O.J., Savedra A.,
Tanuti A., Swanson P.A., Brindeiro R., Schable C., Pieniazek D.,
Rayfield M.;
"HIV-1 subtypes among blood donors from Rio de Janeiro, Brazil.";
Submitted (NOV-1997) to the EMBL/GenBank/DDBJ databases.
EMBL, AF034019; AAC79271.1;
InterPro; IPR000777; GP120.
Pfam: PF00516; GP120;
Pfam: PF00516; GP120;
AAIDS: Coat protein; Glycoprotein.
                                                                                                                                                                                                                                                                                 SEQUENCE FROM N.A.
STRAIN-C-125 / JCM 9153;
MEDLINE-20512582; PubMed-11058132;
MEDLINE-20512582; PubMed-11058132;
Takami H., Nakasone K., Takaki Y., Maeno G., Sasaki R., Masui N., Fuji F., Hirama C., Nakamura Y., Ogasawara N., Kuhara S.,
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                                                                                                                                                                                                      Bacteria; Firmicutes; Bacillus/Clostridium group; Bacillales; Bacillaceae; Bacillus.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      75.6%; Score 34; DB 16; Length 71; 100.0%; Pred. No. 21; tive 0; Mismatches 0; Indels
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115 AA; 12755 MW; 6551E67B32DCF56C CRC64;
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01-MAY-1999 (TrEMBLrel. 10, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
Envelope glycoprotein C2V3 region (Fragment).
                                                                                   01-OCT-2000 (TrEMBLrel. 15, Created)
01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-MAR-2002 (TrEMBLrel. 20, Last annotation update)
Hypothetical protein BH1226.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human immunodeficiency virus type 1.
Viruses; Retroid viruses; Retroviridae; Lentivirus
                                               71 AA.
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Best Local Similarity 100.
Matches 6; Conservative
                                               PRELIMINARY;
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                                                                                                                                                                                     Bacillus halodurans.
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Matches 5; Conservat
                                                                                                                                                                                                                                                  NCBI_TaxID=86665;
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43 NATEYE 48
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SEQUENCE
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                                                                  Q9KDI7;
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Q9YXT3
                           Q9KDI7
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                                                                                                                                           Eukaryota; Metazoa; Nematoda; Chromadorea; Rhabditida; Rhabditoidea;
Rhabditidae; Peloderinae; Caenorhabditis.
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J. Bacteriol. 184:2005-2018(2002).

EMBL. AED10481; AED10481; AL93886.1; -. GEOURICE 628 AA; 73857 FW; 013321316E4829B4 CRC64;
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EMBL: Z50755, 0A890635.1; -.
InterPro: IPR001232; Skpl.
FMART: SM00512: Skpl.
SMART: SM00512: Skpl: 1.
SEQUENCE 182 AA; 21178 MM; 4BE38A19C4FA8124 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           77.8%; Score 35; DB 16; Length 62
62.5%; Pred. No. 1.1e+02;
Live 3; Mismatches 0; Indels
                                                                                                                                                                                                                                                  Coles L.;
Submitted (AUG-1995) to the EMBL/GenBank/DDBJ databases.
Q21968;
01-NOV-1996 (TrEMBLrel. 01, Created)
01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        01-JUN-2002 (TrEMBLrel. 21, Created)
01-JUN-2002 (TrEMBLrel. 21, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
Terratricopeptide repeat family protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Fusobacterium nucleatum (subsp. nucleatum).
Bacteria; Fusobacteria; Fusobacterium.
NCBI_TaxID=76856;
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SEQUENCE FROM N.A.
STRAIN=ATCC 25586;
MEDLINE-21866394;
PUDMed=11889109;
                                                                                                                                                                                                                                                                                                    SEQUENCE FROM N.A.
MEDLINE=99069613; PubMed=9851916;
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Matches 6; Conservative
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                                                                                                  R12H7.3. Caenorhabditis elegans.
                                                                                                                                                                                                                           SEQUENCE FROM N.A.
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161 NNATLFFK 168
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127 NNASYYK 134
                                                                                   R12H7.3 protein.
                                                                                                                                                                                 NCBI_TaxID=6239;
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Q8RI47; **Q8RI47** 

RESULT 3 Q8RI47

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Matches

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Gaps

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MEDLINE-21128732; PubMed=11234002;
Cole S.T., Eiglmeier K., Parkhill J., James K.D., Thomson N.R., Wheeler P.R., Honore N., Garnier T., Churcher C., Harris D., Mungall K., Basham D., Brown D., Chillingworth T., Connor R., Davies R.M., Devlin K., Duthoy S., Feltwell T., Fraer A., Hamlin N., Holroyd S., Hornsby T., Jagels K., Lacrolx C., Maclean J., Moule S., Murphy L., Oliver K., Quail M.A., Rajandream M.A., Rutherford K.M., Rutter S., Seeger K., Simon S., Simmonds M., Skelton J., Squares R., Barrell B.G.;
                                                                                                               SEQUENCE FROM N.A.
STRAIN-CV. CHIHOKU;
MEDLINE-97393494; PubMed-9249988;
MEDLINE-97393494; PubMed-9249988;
MEDLINE-97393494; DubMed-9249988;
MACCUMLA M., Takezawa D., Arakawa K., Yoshida S.;
"Accumulation of 19-Kna plasma membrane polypeptide during induction of freezing tolerance in wheat suspension-cultured cells by abscisic
  Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Pooldeae;
Triticeae; Triticum.
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NCBI_TaxID=1769;
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                                                                                                                                                                                                                                                                              Plant Cell Physiol. 38:707-716(1997).
EMBL; U80037; AAB38504.1; -.
SEQUENCE 182 AA; 19009 MW; 799B9994AEA87DCB CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Hypothetical protein, Complete proteome.
SEQUENCE 273 AA; 30831 MW; D5B765A65EF39549 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-JUN-2001 (TrEMBLrel. 17, Created)
01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
01-OCT-2001 (TrEMBLrel. 18, Last annotation update)
Hypothetical protein ML0126.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      01-JUN-2001 (TrEMBLrel. 17, Created)
01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              "Massive gene decay in the leprosy bacillus.";
Nature 409:1007-1011(2001).
EMBL; AL583917; CAC29634.1; -.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Ź
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Best Local Similarity
6; Conserve
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                                                                      NCBI_TaxID=4565;
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20 NNAQEYE 26
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Q9CJ05;
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Chen L.J., Chai Y.J., Chen P.W.;
Chen L.J., Chai Y.J., Chen P.W.;
A rice embryo-specific gene with high homology to soybean GmPM3 gene,
a rice embryo-specific gene.";
Submitted (MAY-1996) to the EMBL/GenBank/DDBJ databases.
EMBL; U57639; AAD10377.1;
SEQUENCE 173 AA; 18287 MW; 63FAZF778BB7259D CRC64;
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Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Pooldeae;
Triticeae; Hordeum.
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MOTIS P.C., Ranford J.C.;
MOTIS P.C., Ranford J.C.;
"Hordeum vulgare hydrophobic embryo-associated protein PM19 mRNA.";
Submitted (DEC-1999) to the EMBL/GenBank/DDBJ databases.
EMBL; AR218627; AAR29532.1; -.
EMBL; AR218627; AAR29532.1; -.
                                                                 OJZREB;
01-MAY-1999 (TrEMBLrel. 10, Created)
01-MAY-1999 (TrEMBLrel. 10, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
Hydrophobic LEA-like protein.
Oryza sativa (Rice).
Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnollophyta; Liliopsida; Poales; Poaceae;
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Pred. No. 52;
0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-0CT-2000 (TrEMBLrel. 15, Last annotation update)
Plasma membrane associated protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Last sequence update)
Last annotation update)
                                                173 AA.
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P93615;
01-MAY-1997 (TEMBLEEL. 03, Created)
01-MAY-1997 (TEMBLEEL. 03, Last sequen
01-DEC-2001 (TEMBLEEL. 19, Last annota
ABA induced plasma membrane protein PM
WTABAAPM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PRT;
                                                PRT;
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85.7%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Hordeum vulgare (Barley).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   6; Conservative
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                                             PRELIMINARY;
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Best Local Similarity
Matches 6; Conserv
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                                                                                                                                                                                                                                                                                   NCBI_TaxID=4530;
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47 NGATEYE 53
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47 NGATEYE 53
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         NCBI_TaxID=4513
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                                             Q9ZRF8
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P93615
RESULT 6
092RF8
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Q9M625
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Q91655
Q91655;
Q1-NOV-1996 (TrEMBLrel. 01, C.
01-NOV-1996 (TrEMBLrel. 01, L.
01-NOV-1996 (TrEMBLrel. 00, L.
Gene 17 protein.
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Best Local Similarity 62.5°,
The 5; Conservative
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Best Local Similarity 62.5
Matches 5; Conservative
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                                               SEQUENCE FROM N.A.
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318 NSATFYYR 325
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318 NSATEYYR 325
 Glycoprotein VP7.
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091E87
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Q91655
ID Q9165
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                     Lactococcus lactis (subsp. lactis) (Streptococcus lactis).
Bacteria; Firmloutes; Bacillus/Clostridium group; Lactobacillales;
Streptococcaceae; Lactococcus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Ciarlet M., Reggeti F., Pina C.I., Liprandi F.; "Equine rotaviruses with G14 serotype specificity circulate among
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                                                                                                                                                                                                                     75.6%; Score 34; DB 16; Length 319; 85.7%; Pred. No. 89; 1; Indels ive 0; Mismatches 1; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Indels
                                                                                                                                                                     InterProj IPR001173; Glycos_transf_2.
Pfan; PR00535; Glycos_transf_2; 1.
Transferase; Complete protection.
SEQUENCE 319 AA; 37340 WW; DIDAC78AB0A950CA CRC64;
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Last sequence update)
Last annotation update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
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01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-NAX-2002 (TrEMBLrel. 20, Last annotation update)
61ycoprotein VP?
                                                                                                                                                                                                                                                                                                                                                                                             Viruses; dsRNA viruses; Reoviridae; Rotavirus.
                                                                                                                                                                                                                                                                                                                             326 AA
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                                                                                            MEDLINE-21235186; PubMed-11337471;
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MEDLINE=95113988; PubMed=7814511;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     venezuelan horses.";
J. Clin. Microbiol. 32:2609-26:
EMBL: U05348; AAA81914.1;
Interpro: IPRO1963; VP7.
Pfam: PF00434; VP7; 1.
Probom; PD000191; VP7; 1.
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Best Local Similarity 62.5.
--- 5; Conservative
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         Rhamnosyltransferase.
                                                                         SEQUENCE FROM N.A.
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318 NSATFYYR 325
                                                                                                                                                                                                                                                                                                                                                                                    Equine rotavirus.
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80 NNADFYF 86
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                                                                                   STRAIN-IL1403
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01-NOV-1996
01-MAR-2002
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Q66772
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Griffin D.D., Nakagomi T., Hoshino Y., Nakagomi O., Kirkwood C.D.,
Parashar U.D., Glass R.I., Gentsch J.R.;
Characterization of nontypeable rotavirus strains from the United
States:Identification of a new rotavirus reasortant (P2A[6],G12) and
rare P3[9] strains related to bowine rotaviruses.";
Submitted (APR-2001) to the EMBL/GenBank/DDBJ databases.
BMBL, AJ311731, ...
InterPro; IPR001963; VP7.
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                                                                                                                                                                           Ciarlet M., Reggeti F., Pina C.I., Liprandi F.; "Equine rotaviruses with G14 serotype specificity circulate among venezuelan horses.";
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62.5%; Pred. No. 91;
ive 3; Mismatches 0; Indels
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                                                                                                                                                                                                                                                                                                                           Pfam; PF00434; VP7; 1.
Probom; PD000191; VP7; 1.
SEQUENCE 326 AA; 37352 MW; 9095E64B13933E29 CRC64;
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Prodom: PD00191; VP7; 1.
PROSITE; PS00343; GRAM POS_ANCHORING; UNKNOWN_1
SEQUENCE 326 AA; 37119 MW; 8B97ED1DBBDLC981 CRC64;
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01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-MAR-2002 (TrEMBLrel. 20, Last annotation update)
Capsid protein.
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Last annotation update)
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62.5%; Pred. No. 91;
tive 3; Mismatches 0;
Equine rotavirus.
Viruses; dsRNA viruses; Recviridae; Rotavirus.
NCBI_TaxID=10937;
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NCBL_TaxID=10941;
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                                                                                                                                                                                                                                                     J. Clin. Microbiol. 32:2609-2612(1994).
EMBL; U05349; AAA67342.1; -.
InterPro; IPR001963; VP7.
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Bukaryota; Viridiplantae; Chlorophyta; Pedinophyceae; Pedinomonadales;
Pedinomonadaceae; Pedinomonas.

NCBI_TaxID=3159;
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                                                                                                         SEQUENCE FROM N.A.
MEDLINE=96312514; PubMed=8700860;
Brown D.D., Wang Z., Furlow J.D., Kanamori A., Schwartzman R.A.,
Prown D.F., Phider A., Purlow J.D., Kanamori A., Schwartzman R.A.,
"The thyroid hormone-induced tail resorption program during Xenopus laevis metamorphosis.";
Xenopus laevis (African clawed frog).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Amphibia; Batrachia; Anura; Mesobatrachia; Pipoidea; Pipidae;
Xenopodinae; Xenopus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Turmel M., Lemieux C., Burger G., Lang B.F., Otis C., Plante I.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Submitted (DEC-1998) to the EMBL/GenBank/DDBJ databases.
-!- CATALYTIC ACTIVITY: NADH + UBIQUINONE = NAD(+) + UBIQUINOL.
EMBL; AF116775; AAD19671.1; -.
InterPro: IPRO001750; Oxidored_q1.
Pfam: PF00361; oxidored_q1: 1.
Mitochondrion; NAD; Oxidoreductase; Ubiquinone.
SEQUENCE 440 AA; 51380 MW; C6D6CBAD72549B4C CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            h 75.6%; Score 34; DB 8; Length 440; Similarity 85.7%; Pred. No. 1.2e+02; 6; Conservative 0; Mismatches 1; Indels
                                                                                                                                                                                                      EMBL; U41860; AAC59876.1; -.
EMBL; U41860; AAC59876.1; -.
InterPro; IPR001507; Endoglin/CD105.
Pfam: PF00100; zona_pellucida; 1.
SEQUENCE 357 AA; 39090 MW; 5493352C8BEA2IE6 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  01-WAY-1999 (TrEMBLrel. 10, Created)
01-MAY-1999 (TrEMBLrel. 10, Last sequence update)
01-MAY-2002 (TrEMBLrel. 20, Last annotation update)
NADH dehydrogenase subunit 2 (EC 1.6.5.3).
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STRAIN-UTEX LB 1350;
MEDLINE-99418884; PubMed-10488238;
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STRAIN-UTEX LB 1350;
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Best Local Similarity
Matches 6; Conserv
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238 NNALFYF 244
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Gapop 10.0 , Gapext 0.5
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## /SIDSI/godata/geneseq/geneseqp-embl/AA1991.DAT:\*/SIDSI/gogdata/geneseq/geneseqp-embl/AA1992.DAT:\*/SIDSI/gogdata/geneseq/geneseqp-embl/AA1993.DAT:\*/SIDSI/gogdata/geneseq/geneseqp-embl/AA1993.DAT:\*/SIDSI/gogdata/geneseq/geneseqp-embl/AA1994.DAT:\* /SIDS1/gcgdata/geneseq/geneseqp-emb1/AA1999.DAT:\* /SIDS1/gcgdata/geneseq/geneseqp-emb1/AA2000.DAT:\* /SIDS1/gcgdata/geneseq/geneseqp-emb1/AA2001.DAT:\* /SIDS1/gcgdata/geneseq/geneseqp-emb1/AA2001.DAT:\* /SIDS1/gcgdata/geneseq/geneseqp-embl/AA1995.DAT:\*/SIDS1/gcgdata/geneseq/geneseqp-embl/AA1996.DAT:\*/SIDS1/gcgdata/geneseq/geneseqp-embl/AA1997.DAT:\* /SIDS1/gcgdata/geneseq/genesegp-embl/AA1998.DAT:\*

1: /SIDS1/gggdata/geneseq/geneseqp-emb1/AA1980.DAT:\*
2: /SIDS1/gggdata/geneseq/geneseqp-emb1/AA1981.DAT:\*
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7: /SIDS1/gggdata/geneseq/geneseqp-emb1/AA1986.DAT:\*
8: /SIDS1/gggdata/geneseqf/geneseqp-emb1/AA1987.DAT:\*
9: /SIDS1/gggdata/geneseqf/geneseqp-emb1/AA1989.DAT:\*
10: /SIDS1/gggdata/geneseqf-emb2/AA1989.DAT:\*
11: /SIDS1/gggdata/geneseqf-emb2/AA1989.DAT:\*
12: /SIDS1/gggdata/geneseqf-emb1/AA1989.DAT:\*
13: /SIDS1/gggdata/geneseqf-emb1/AA1989.DAT:\*
14: /SIDS1/gggdata/geneseqf-emb1/AA1989.DAT:\*

A\_Geneseq\_101002:\*

Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Description	Anti-angiogenic D3	Anti-anglogenic D3	Domaine 3, bradyki	Human kininogen D3	Human high mol.wt.	Novel human diagno	Novel human diagno	Human high molecul	Novel human diagno	Human high molecul
SUMMARIES	AAY95410	AAY95408	AAR33350	AAB37447	AAY95426	ABG21102	ABG21099	ABB78707	ABG21101	ABB78710
DB	21	21	14	21	21	22	22	23	22	23
% Query Match Length DB	16	32	117	122	123	248	369	626	644	644
% Query Match	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0
Score	84	84	84	84	84	84	84	84	84	84
Result No.	п	7	e	4	ഗ	9	7	æ	6	10

	11	<b>ھ</b> د	CI C	26	18	AAW54336	Bradykinin analogo
	13	) ()	$^{\circ}$	434	3 2	AAP40633	Bradykinin protein
	14 15	- 19	<b>~</b> 9	16 235	23	AABU8553 ABG60077	Peptide identified Human DITHP polype
	16 17	40	4 8	436	21	AAP40257 AAB37455	Bradykinin protein Human kininogen D3
	18 19	45	53.6	10	21 21	AAY95405 AAY95409	Anti-angiogenic D3 Anti-angiogenic D3
	20	<u>د</u> د	ω.	554	23	ABP27632	Streptococcus poly
•	22	4 4	M (A)	875	200	AAY34308 AAY34381	Porphorymonas ging Porphorymonas ging
	23	m <del></del>	ᇽ	466	23	ABB48863	Listeria monocytog
:	25	0	001	966	533	ABB91160	Herbicidally activ
	26	00	~ ~	152	22	AA006226 AAG32794	Human polypeptide
	28	. 0	1	153	21	AAG32793	Zea mays protein f
	30	00	~ ~	166	21	AAG32792	Zea mays protein f
	31		-	1191	55	ABB60775	Drosophila melanog
	32	o 0	ωu	12	21	AAY95407	Anti-angiogenic pe
	4.6	, œ	വ	404	23	AAU89265	Insulin/insulin-li
	35		· CO I	42	22	AAG74378	Human colon cancer
	3.0	œ c	ഥവ	44	53	AAU90767	Insulin/insulin-li
	38	0 00	വര	128	21	AAY81189	Human mutant cysta Human mutant cysta
	39	· &	2	156	18	AAW20537	H. pylori cytoplas
	0 7	ω (	ro r	222	21	AAB52576	Helicobacter pylor
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RES	ESULT 1						
AAY9:	5410 AAY95	410 star	tandard;	Peptide	je;	16 AA.	
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X X	AAY954	410;					
ដ	. 25-SEP	-2000	(first	t entry	2		
X C	Ant 1-a	-angingenic	nic D3	pentid	ą		
×	5	55045:	24	i dod	;		
K K	Anti-angiogenic endothelial cel	ngiogen elial (	nic; ar	olifer	rati	inhibitor; apoptosis;	ън
KW KW	rheumatoid therapy; hu	toid art y; human	arthriti man; D3	ritis; cytostati D3 peptide.	tost de.	c, antiarthr	antirheumatic;
XX OS	ното в	sapiens					
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PF	02-DEC	-1999;	09WC	9WO-US2846	165.		
PR	16-DEC	-1998;	9808	5-01124	127.		
X & a	(UTEM )	UNIV TE	TEMPLE.	61			
χX	(שררע)		4				
PI	McCrae	RK;					
DR	WPI; 2	2000-44;	42247/38	ë.			
PT	Composition		for in	inhibiting		angiogenesis and	endothelial cell

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                                                                                         The present sequence is that of a D3 peptide derived from human him of wt. Kininogen (HK) domain 3 (see AAX95426). The D3 peptide inhibits endothelial cell proliferation and thus possesses anti-anglogenic activity. It is an example of D3 peptides of the invention (see AAY95405-256) that are analogues of certain sites in the HK domain 3, in this case amino acid residues Asn275-Lys282. The PEPtides inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting andiogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell proliferation of the retina are treated. The Internation of the retina are treated. The Internation of fibroblast growth factor-induced HUVEC cell
proliferation, inducing endothelial cell apoptosis and treating cancer, rheumatoid arthritis, and ocular disorders comprises a kininogen domain
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100.0%; Pred. No. 1.4e-07;
ive 0; Mismatches 0;
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                                                           Claim 7; Page 26; 44pp; English
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les 16; Conservative
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                                                                                                                                                                                                                                                                                                                      proliferation.
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                                 3 analog
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The present sequence is that of a D3 peptide derived from human

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Domain 3; human; kininogen; heavy chain; low molecular weight; plasma; trypsin; platelet; activation; granule contents; hemostasis; thrombin; tissue plasminogen activator;thrombosis; inflammatory response; endothelial cell; vow Willebrand factor;
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high mol.wt. kininogen (HK) domain 3 (see AAY95426). The D3 peptide inhibits endothelial cell proliferation and thus possesses anti-angiogenic activity. It is an example of D3 peptides of the invention (see AAY95465-26) that are analogues of certain sites in the HK domain 3, in this case amino acid residues Asn275-Lys282. The peptides inhibit endothelial cell proliferation and may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting angiogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell proliferation, and inducing endothelial cell apoptosis. Cancer, rheumatchid arthritis, and ocular disorders cell apoptosis. Cancer, rheumatchid arthritis, and ocular disorders
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/note= "1
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les 16; Conserv
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Peptide
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Composition for inhibiting angiogenesis and endothelial cell proliferation, inducing endothelial cell apoptosis and treating cancer, rheumatoid arthritis, and ocular disorders comprises a kininogen domain
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   The present sequence is that of domain 3 of human high mol.wt. kininogen (HK). The invention provides peptides (see AAY95405-24) that are analogues of certain sites in the HK domain 3, specifically Asn275-Lys282, Cys246-Cys249, Leu331-Tyr338 and Tyr299-Ser314. The peptides, in which native Cys residues may be replaced by Ala residues, in which native Cys residues may be nod may also induce endothelial cell apoptosis. Compositions including the peptides are used in claimed methods for inhibiting angiogenesis, inhibiting endothelial cell proliferation, and inducing endothelial cell apoptosis. Cancer, rheumatoid arthritis, and ocular disorders characterized by undesired vascularization of the retina are treated.
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     AAY95426 standard; Peptide; 123
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Matches 16; Conserv
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                                        AAY95426;
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Domain 3 also inhibits endothelial cell activation shown by a decrease in secretion of endothelial cell contents such as tissue plasminogen activator and von Willebrand factor. Domain 3 functions to inhibit cell activation by blocking thrombin binding to its target cells, the peptide is a selective inhibitor of thrombin-induced platlet
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Inhibition of mammalian legumain or legumain-related endopeptidase by cystatin involves interaction with second papain-non-reactive site of cystatin
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                                                                                                                                                                      Length 117;
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100.0%; Pred. No. 1.3e-06;
11ve 0; M1smatches 0;
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                                                                                                                                                                                                                                                                    30 NNATFYFKIDNVKKAR 45
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                                                                                                                                                                                        Local Similarity
nes 16; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Human kininogen D3
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                                                                                                                                  117 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WO200064945-A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Homo sapiens
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       02-NOV-2000.
                                                                                             activation.
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                                                                                                                                  Sequence
                                                                                                                                                                    Query Match
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Length 123; Indels Tang YT;

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New isolated polynucleotide and encoded polypeptides, useful in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
                                                                                                                                                                                                                                                                                                             Claim 20; SEQ ID No 51458; 103pp; English.
                                                                      30-MAR-2001; 2001WO-US08631.
                                                                                                  2000US-0540217
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                                                                                                                                                                      Drmanac RT, Liu C,
                                                                                                                                                                                                 WPI; 2001-639362/73.
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Matches 16; Conserv
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                                                                                                                                           (HYSE-) HYSEQ INC
                                                                                                                                                                                                              N-PSDB; AAS85286
                WO200175067-A2.
                                                                                                  31-MAR-2000;
                                                                                                                23-AUG-2000;
                                                                                                                                                                                                                                                                                    biodiversity
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                                          11-0CT-2001
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Note: The sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/published_pct_sequences.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          and to produce other types of data and products dependent on DNA and amino acid sequences. ABG00010-ABG30377 represent novel human diagnostic amino acid sequences of the invention.
                                                                                                                                                                                                                                                                 New isolated polynucleotide and encoded polypeptides, useful in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human; chromosome mapping; gene mapping; gene therapy; forensic; food supplement; medical imaging; diagnostic; genetic disorder.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        100.0%; Score 84; DB 22; Length 248; 100.0%; Pred. No. 2.9e-06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    0; Indels
                                                                                                                                                                                                                                                                                                                                                                  The invention relates to isolated polynucleotide (I) and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Mismatches
                                                                                                                                                                                                                                                                                                                                   Claim 20; SEQ ID No 51461; 103pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Novel human diagnostic protein #21090.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABG21099 standard; Protein; 369 AA.
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                                                                                               30-MAR-2001; 2001WO-US08631.
                                                                                                                           31-MAR-2000; 2000US-0540217.
23-AUG-2000; 2000US-0649167.
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                                                                                                                                                                                                                          WPI; 2001-639362/73
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             248 AA;
                                                                                                                                                                    (HYSE-) HYSEQ INC
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                                       WO200175067-A2.
              Homo sapiens.
                                                                                                                                                                                                                                                                                                             biodiversity
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                                                                    11-OCT-2001
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                                                                                                                                                                                                                                     for identifying expressed genes. (I) is useful in gene therapy techniques to restore normal activity of (II) or to treat disease states involving (II). (II) is useful for generating antibodies against it, detecting or quantitating a polypeptide in tissue, as molecular weight markers and as a food supplement. (II) and its binding partners are useful in medical imaging of sites expressing (II). (I) and (II) are useful for treating disorders involving aberrant protein expression or biological activity. The polypeptide and polynucleotide sequences have applications in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits to assess biodiversity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      and to produce other types of data and products dependent on DNA and amino acid sequences. ABG00010-ABG30377 represent novel human diagnostic amino acid sequences of the invention.

Note: The sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from WIPO
The invention relates to isolated polynucleotide (I) and polypeptide (II) sequences. (I) is useful as hybridisation probes, polymerase chain reaction (PCR) primers, oligomers, and for chromosome and gene mapping, and in recombinant production of (II). The polynucleotides are also used in diagnostics as expressed sequence tags
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    100.0%; Score 84; DB 22; Length 369; 100.0%; Pred. No. 4.5e-06;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ftp.wipo.int/pub/published_pct_sequences.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Pred. No. 4.5
0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   333 NNATEYEKIDNVKKAR 348
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polypeptide (II) sequences (I) is useful as hybridisation probes, polypeptide (II) sequences (I) by useful as hybridisation probes, polymerase chain reaction (PCR) primers, oligomers, and for chromosome and gene mapping, and in recombinant production of (II). The polymerese chain reaction (PCR) primers, oligomers, and for chromosome polymered are also used in disgnostics as expressed sequence tags for identifying expressed genes, (I) is useful in gene therapy techniques to restore normal activity of (II) or to tread disease states involving (II). (II) its useful for generating antibodies against it, detecting or quantitating a polypeptide in tissue, as molecular weight markers and as a food supplement. (II) and (II) and (II) are useful for treating disorders involving aberrant protein expression or biological activity. The polypeptide and polymericide sequences have applications in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits to assess biodiversity amino acid sequences. Absoloulo Absolution.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              D5 receptor; angiogenesis; endothelial cell; cytostatic; antitumour; antiatherosclerotic; vasotropic; vulnerary; tranquilliser; thrombolytic;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Note: The sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from WIPO
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     New isolated polynucleotide and encoded polypeptides, useful in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human; kininogen; high molecular weight kininogen; HK; D5 domain;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ö
food supplement; medical imaging; diagnostic; genetic disorder
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            invention relates to isolated polynucleotide (I) and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human high molecular weight kininogen (HK) protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       at ftp.wipo.int/pub/published_pct_sequences
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Claim 20; SEQ ID No 51460; 103pp; English.
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                                                                                                                                                                                                                                                                                                                                                                                             Tang YT;
                                                                                                                                                                                                         30-MAR-2001; 2001WO-US08631.
                                                                                                                                                                                                                                                          31-MAR-2000; 2000US-0540217. 23-AUG-2000; 2000US-0649167.
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                              WPI; 2001-639362/73
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          644 AA;
                                                                                                                                                                                                                                                                                                                                         HYSE-) HYSEQ INC.
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                                                                                                    WO200175067-A2.
                                                      Homo sapiens.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         biodiversity
                                                                                                                                                     11-0CT-2001.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         The present invention describes an isolated polypeptide (I) that

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of human kininogen, or biologically active

corresponds to the D5 domain of human kininogen, or biologically active

corresponds and correct the correct of the biological or activates signalling pathways leading to the

correction of apoptosis in EC; and/or (d) inhibits the signalling pathway required for maintenance of EC viability. (I) has cytostatic,

correction, antiantherosciencic, vasotropic, vulnerary, tranquilliser,

thrombolytic, ophthalmological, gynaecological, antiulcer, antidiabetic,

correction antianthological, gynaecological, antiulcer, antidiabetic,

correction antibody (IX) specific for an epitope of (I) is useful for inhibiting

correction antibody (IX) specific for an epitope of (I) is useful for inhibiting

correction antibody (IX) specific for an epitope of (I), a D5 fusion polypeptide

correction antibody (IX) specific for an epitope of (I), a D5 fusion polypeptide

correction and adjustion, proliferation, invasion, or angiogenesis, or

composition (X) comprising (I), (II), can be used for treating

correction proliferation, invasion or angiogenesis. (I), (II), or (III)

correction associated with undesired EC

composition, proliferation, invasion or angiogenesis. (I), (II), or (III)

correction a used for isolating or enriching cells expressing D5 domain binding

correction acell mixture. The present sequence represents the mature

correction of the present invention invention in the exemplification of the present invention.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Novel human kininogen D5 domain polypeptides useful for treating conditions associated with endothelial cell migration, proliferation, invasion or anglogenesis, e.g. arthritis, macular degeneration, benign
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     the exemplification of the present invention
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                                              Location/Qualifiers
384..508
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                                                                                    71abel D5_domain
                                                                                                                                                                                                                                                     24-JUL-2001; 2001WO-US23185
                                                                                                                                                                                                                                                                                                           24-JUL-2000; 2000US-220194P
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Local Similarity 100.
nes 16; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                fazar AP, Juarez JC;
                                                                                                                                                                                                                                                                                                                                                               (ATTE-) ATTENUON LLC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 WPI; 2002-393611/42.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      626 AA;
                                                                                                                                                   WO200214369-A2
Homo saptens
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                                                                         Domain
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The present invention describes an isolated polypeptide (1) that
corresponds to the D5 domain of human kininogen, or biologically active
peptide fragment, homologue or functional derivative, and which:
(a) inhibits angiogenesis; (b) binds to the D5 binding site on
endothelial cells (EC); (c) activates signalling pathways leading to the
introduction of apoptosis in EC; and/or (d) inhibits the signalling
introduction of apoptosis in EC; and/or (d) inhibits the signalling
of antitumour, antiatherosclerotic, vasotropic, vulnerary, tranquilliser,
thrombolytic, ophthalmological, gynaecological, antiulcer, antidiabetic,
antiarthritic, antiandiogenic, antiapoptotic and endocrine activities.
An antibody (IX) specific for an epitope of (1) is useful for inhibiting
tumour growth or angiogenesis in a subject. (1), a D5 fusion polypeptide
(II) or a diameric or trimeric fusion polypeptide (III) can be used for
inhibiting EC apoptosis. An angiogenic EC-targeting pharmaceutical
composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (X) composition (
ophthalmological; gynaecological; antiulcer; antidiabetic; antiarthritic; antianglogenic; antiapoptotic; endocrine; apoptosis; gene therapy.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         domain binding
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Novel human kininogen D5 domain polypeptides useful for treating conditions associated with endothelial cell migration, prolliferation, invasion or anglogenesis, e.g. arthritis, macular degeneration, benign
                                                                                                                                                                                        9..644
|Tabol= mature_human_high_molecular_weight_kininogen
18..614
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Disclosure; Fig 1B-E; 84pp; English.
                                                                                                                                  Socation/Qualiflers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   /label- D5_domain
                                                                                                                                                                ...18
'label= signal
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          24-JUL-2000; 2000US-220194P.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Novel human kininogen
conditions associated
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Mazar AP, Juarez JC;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (ATTE-) ATTENUON LLC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WPI; 2002-393611/42.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      644 AA;
                                                                                                                                                                                                                                                                                                                                                                   Disulfide-bond
Disulfide-bond
Disulfide-bond
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Disulfide-bond
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Disulfide-bond
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                                                                                                                                                                                                                                                                                                                                                Disulfide-bond
                                                                                    Homo sapiens
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                                                                                                                                                          Peptide
                                                                                                                                                                                                                   Protein
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thrombin-induced platelet; prevention; platelet aggregation;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Administration of a peptide or multimer related to bradykinin or other
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          disclosed peptides and multimers can be used for the inhibition of thrombin-induced platelets or other cells. They can also be used for preventing platelet aggregation, or inhibiting ADP-induced activation. This is useful to prevent arterial occlusions arising from coronary thrombosis and stroke.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         activation and aggregation - by admin. of new
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human; chromosome mapping; gene mapping; gene therapy; forensic; food supplement; medical imaging; diagnostic; genetic disorder.
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2.5e-06;
hes 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 92.9%; Score 78; 100.0%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Novel human diagnostic protein #21096.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Disclosure; Page 44; 73pp; English.
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                                                                                                                                                                                                        AAW54336 standard; peptide; 26 AA.
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100.08; Pr
0;
                                                                                                                                                                                                                                                                                                                                                                                  Bradykinin analogous peptide 19.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Inhibition of platelet actival or known bradykinin analogues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               96WO-US09940
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     950s-0000096
                                                                                                                                                                                                                                                                                                                            (first entry)
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                                    293 NNATFYFKIDNVKKAR 308
1 NNATEYFKIDNVKKAR 16
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Matches 15; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Hasan AAK, Schmaier AH;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   1 NATEYEKIDNVKKAR 15
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                           activation.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           26 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           WO9641640-A1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            07-JUN-1996;
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                                                                                                                                                                                                                                                                                                                                                                                                                                         Inhibition;
ADP-induced
                                                                                                                                                                                                                                                                                                                         30-JUL-1998
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Synthetic.
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Length 644; Indels

100.0%; Score 84; DB 23; 100.0%; Pred. No. 8.3e-06; ive 0; Mismatches 0;

Conservative

Local Similarity nes 16; Conserv

Best Loc Matches

Query Match

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20-DEC-2000 (first entry)
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Matches 11; Conservative
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290 HDGTFYFKIDTVKKA 304
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                                                                                                                                                                                                                    WPI; 1984-216122/35.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           434 AA;
                                                                                                                                                                                                                                           N-PSDB; AAN40314.
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JP59125896-A.
                                                                                    07-JAN-1983;
                                                                                                                             07-JAN-1983;
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                                           20-JUL-1984.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          31-AUG-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAB08553;
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NAME OF STATES O
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             à
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 The invention relates to isolated polynucleotide (I) and polypeptide (II) sequences. (I) is useful as hybridisation probes, polymerase chain reaction (PGR) primers, oligomers, and for chromosome and gene mapping, and in recombinant production of (II). The compliant production of (II). The polynucleotides are also used in diagnostics as expressed sequence tags for identifying expressed genes. (I) is useful in gene therapy techniques to restore normal activity of (II) or to treat disease states involving (II). (II) is useful for generating antibodies against it, detecting or quantitating a polypeptide in tissue, as molecular weight markers and as a food supplement. (II) and its binding partners are useful in medical ingoing of sites expressing (II). (I) and (II) are useful in medical ingoraters involving aberrant protein expression or biological activity. The polypeptide and polynucleotide sequences have applications in diagnostics, forensics, gene mapping, identification of mutations and to produce other types of data and products dependent on DNA and and can demonstrate and sequences of the invention.

The sequence data for this patent did not appear in the printed specification, but was obtained in electronic format directly from WIPO at the invitor. In the printed specification, but was obtained in electronic format directly from WIPO at the invitor.
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                                                                                                                                                                                                                                                                                                                           New isolated polynucleotide and encoded polypeptides, useful in diagnostics, forensics, gene mapping, identification of mutations responsible for genetic disorders or other traits and to assess biodiversity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 435;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 70; DB 22; Length 43
Pred. No. 0.0013;
0; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Bradykinin protein precursor: type II (pKG146, pKG254).
                                                                                                                                                                                                                                                                                                                                                                                                                                      Claim 20; SEQ ID No 51464; 103pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAP40633 standard; Protein; 434 AA.
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/label- bradykinin
391..395
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Blood pressure; kininogen; probe..
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               83.3%;
87.5%;
                                                           30-MAR-2001; 2001WO-US08631.
                                                                                                    31-MAR-2000; 2000US-0540217.
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Matches 14; Conservative
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                                                                                                                                                                                                                  Drmanac RT, Liu C,
                                                                                                                                                                                                                                                           WPI; 2001-639362/73.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        435 AA;
                                                                                                                                                                      HYSE-) HYSEQ INC
                                                                                                                                                                                                                                                                                  N-PSDB; AAS85292
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                  11-0CT-2001
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Bradykinin is a peptide consisting of nine amino acids. It has the biological effect of decreasing blood pressure. Although kininogen is known as a protein-precursor of bradykinin, its structure is unknown because of the difficulty in collecting large enough samples of kininogen for structural investigation.
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                                                                                                                                                                                                                                                                   c-Dna fragment of protein precursor - used to code bradykinin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 434;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 59; DB 5;
Pred. No. 0.091;
2; Mismatches
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                                                                                                                                                                                                                                                                                                                                 Disclosure; Fig 2; 6 pp; Japanese.
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                                                                                                                     (MITU ) MITSUBISHI CHEM IND KK.
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                                                             83JP-0000984.
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83JP-0000984
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Human, DITHP; diagnostic and therapeutic polypeptide; bone; testis; skin; cell proliferative disorder; cancer; tumour; autoimmune disorder; brain; inflammatory disorder; viral infection; bacterial infection; seizure; fungal infection; parasitic infections; developmental disorder; breast; endocrine disorder; metabolic disorder; neurological disorder; cervix; gastrointestinal disorder; transport disorder; gene therapy; kidney; adrenal gland; bone marrow; lung; ovary; pancreas; prostate; spieen;
                          The specification describes a method for identifying a peptide having a particular function. The method comprises preparing a database of polynucleotides and polypeptides of unknown functions, screening the database for a combination of nucleotides or amino acids indicative of the peptide with a particular function, and identifying polynucleotides and proteins which contain the peptide. The method is used to identify precursor peptides with an amidated C-terminus, especially polypeptide hormones, for studying physiologically active substances. The present sequence represents a peptide which was identified using the method of
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                                                                                                                                                                                                                  Score 57; DB 21; Length 16;
Pred. No. 0.0051;
0; Mismatches 0; Indels
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                                                                                                                                                                                                          67.9%; Scc.
100.0%; Pre
Claim 16; Page 20; 40pp; French
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human DITHP polypeptide #135.
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Best Local Similarity
                                                                                                                                                                                          16 AA;
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                                                                                                                                                               the invention.
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05-SEP-2000;
05-SEP-2000;
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05-SEP-2000;
06-SEP-2000;
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Search completed: March 13, 2003, 19:12:03 Job time: 43 secs

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The invention relates to human diagnostic and therapeutic (dithp)

Complete and their associated polypeptides (DITHP polypeptides).

The sequences of the invention are used in the treatment and diagnosis of cell proliferative disorders (e.g. atherosclerosis, cirrhosis), cancers cell proliferative disorders (e.g. atherosclerosis, cirrhosis), cancers cell proliferative disorders (e.g. atherosclerosis, cirrhosis), cancers cervix, kidney, lung, ovary, pancreas, prostate, skin, spleen, testis or thymus), autoimmune/inflammatory disorders (e.g. asthma, bronchitis, psoriasis, osteoporosis), viral infections, bacterial infections, fungal infections, parasitic infections, developmental disorders (e.g. anaemia, epilepsy), selzure disorders (e.g. cerebral palsy, splin bifida), endocrine disorders (e.g. thrombosis, aneurysm), metabolic disorders (e.g. obesity, diabetes), neurological disorders (e.g. stroke, anyotrophic lateral sclerosis, multiple sclerosis), gastrointestinal clisorders (e.g. ulcerative colitis, lysinuria) and transport disorders (e.g. myotronic dystrophy, catatonia, peripheral neuropathy). Sequences ABG59943-ABG60220 represent human DITHP polypetides of the invention.
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                                                                                      Dahl CR;
                                                              Chalup MS, Hillman JI
TF, Yap PE, Dahl CR;
B, Roseberry AM;
, Flores V, Daffo A;
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Pred. No. 0.15;
1; Mismatches 3; Indels
                                                           Lincoln SE, Altus CM, Dufour GE, Cha
xu JY, Wright RJ, Gietzen D, LiurFr,
Bradley DL, Robatgi SD, Harris B,
Peralta CH, David MH, Panzer SR, F
                                                                                                                                                  Inman RR:
                                                       Dufour GE,
tzen D, Liu
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                                                                                                                                                  Chang SC,
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73.3%;
                   (INCY-) INCYTE GENOMICS INC.
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Best Local Similarity 73.3
Matches 11; Conservative
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Marwaha R, Chen AJ,
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N-PSDB; ABK71668.
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                                                         Stuart J,
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Patent No. 523394
Patent No. 523394
Sequence 10, Appl
Sequence 10, Appl
Sequence 10, Appl
Sequence 2, Appl
Patent No. 523394
Patent No. 523394
Patent No. 518541
Sequence 2, Appl
Sequence 31, Appl
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Patent No. 5472945
GENERAL INFORMATION:
APPLICANT: Schmaier, Alvin H.
APPLICANT: Jang, Yongping
TITLE OF INVENTION: Modulation of Blood
TITLE OF INVENTION: Pressure and Inhibition of Platelet Activation
TITLE OF INVENTION: with Kininogen Fragment
NUMBER OF SEQUENCES: 2
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match 100.0%; Score 84; DB 1; Length 117; Best Local Similarity 100.0%; Pred. No. 5.3e-07; Matches 16; Conservative 0; Mismatches 0; Indels
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STREET: 1800 Two Penn Center Plaza
CITY: Philadelphia
CITY: Philadelphia
COUNTY: U.S.A.
ZIP: 19102
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 3.50 inch, 720 Kb
COMPUTER: WordPerfect 5.1
COMPUTER: WordPerfect 5.1
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/193,114B
FILING DATE: 9 February 1994
CLASSIFICATION: 514
PRIOR APPLICATION: 514
PRIOR APPLICATION BATA:
APPLICATION NUMBER: U.S. Application
APPLICATION NUMBER: Serial No. 5472945 07/744,545
FILING DATE: 13 AUGUST 1991
ATTORNEY/AGENT INFORMATION:
AND MANDER AND MANDER SERIAL SERIAL AND MANDER AND MANDER SERIAL 
                       ALIGNMENTS
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REFERENCE/DOCKET NUMBER: 6056-137 C11
TELECOMMUNICATION INFORMATION:
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TELEFAX: (215) 568-5549
TELEX: NO. 5472945e
INFORMATION FOR SEQ ID NO: 1:
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peptide
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Sequence 1, Appli
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Sequence 14, Appl
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                                                                                                                                                                                                                                         March 13, 2003, 19:10:57; Search time 15.3333 Seconds (without alignments) 30.702 Million cell updates/sec
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                               GenCore version 5.1.4\_p5\_4578 Copyright (c) 1993 - 2003 Compugen Ltd.
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2: /cgn2_6/ptodata/2/laa/5B_COMB.pep:*
3: /cgn2_6/ptodata/2/laa/6A_COMB.pep:*
4: /cgn2_6/ptodata/2/laa/B_COMB.pep:*
5: /cgn2_6/ptodata/2/laa/PCTUS_COMB.pep:*
6: /cgn2_6/ptodata/2/laa/pcTUS_COMB.pep:*
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US-08-66-014-3

US-08-66-014-3

US-08-679-493A-85

US-09-134-001C-3050

US-09-134-001C-3050

US-09-282-305-14

US-08-136-136-13

US-08-136-136-13

US-08-479-939-21

US-08-483-21

US-09-134-001C-3554

US-09-134-001C-3554

US-08-487-48-4

US-08-726-625-4

US-08-487-948-4

US-08-08-983-516-4
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US-07-940-861-12
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Maximum Match 100%
Listing first 45 summaries
                                                                                                                                                                           protein search, using sw model
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Gapop 10.0 , Gapext 0.5
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Issued_Patents_AA:*
                                                                                                                                                                                                                                                                                                                                                                                      US-09-461-061A-10
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length: 2000000000
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Match Length DB
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Minimum DB Maximum DB

Database

Title: Perfect score:

Sequence:

OM protein

Run on:

Scoring table:

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Gaps

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36.5 36.5 36.5 36

Score

No.

Result

Gaps

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OTHER INFORMATION: Description of Artificial Sequence: Bradykinin; OTHER INFORMATION: analog
US-08-676-242-15
                                                                                                                                                                                                                                                                                                                                     DB 4; Length 26;
1.1e-06;
                                                                                                                                                                                                                                                                                                                                                                                 Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              APPLICANT: Roger K. Maes and Stephen J. Spatz
TITLE OF INVENTION: Recombinant Poxvirus
TITLE OF INVENTION: Vaccine Against
TITLE OF INVENTION: Feline Rhinotracheitis
NUMBER OF SEQUENCES: 11
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ZIP: 48864
COMPUTER READABLE FORM:
MEDIUM TYPE: Diskette, 5.25 inch, 360 Kb
MEDIUM TYPE: storage
                                                                                                                                                                                                                                                                                                                                                                              0; Mismatches
                  EARLIER FILING DATE: 1995-06-09
EARLIER APPLICATION NUMBER: PCT/US96/09940
EARLIER FILING DATE: 1996-06-07
NUMBER OF SEQ ID NOS: 24
SOFTWARE: Patentin Ver. 2.1
SEQ ID NO 15
LENGTH: 26
                                                                                                                                                                                                                                                                                                                                Query Match 92.9%; Score 78; Best Local Similarity 100.0%; Pred. No. Matches 15; Conservative 0; Mismatch
EARLIER APPLICATION NUMBER: 60/000,096
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               OPERATING SYSTEM: MS-DOS
SOFTWARE: WORDDEFIECT 5.1
SOFTWARE: APPLICATION DATA:
APPLICATION NUMBER: US/08/911,321
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/096,183
FILING DATE: July 26, 1993
ATTORNEY AGENT INFORMATION:
NAME: Jan C. McLeod
REGISTRATION NUMBER: 20,931
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              REFERENCE/DOCKET NUMBER: MSU 4.
TELECOMMUNICATION INFORMATION:
TELEPAX: (517) 347-4100
TELEFAX: (517) 347-4103
TELEX: NO. 6010703e
INFORMATION FOR SEQ ID NO: 3:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADDRESSEE: Ian C. McLeod
STREET: 2190 Commons Parkway
CITY: Okemos
STATE: Michigan
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ; Sequence 3, Application US/08911321
; Patent No. 6010703
; GENERAL INFORMATION:
                                                                                                                                                                         TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        COMPUTER: IBM Compatible OPERATING SYSTEM: MS-DOS
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                                                                                                                                                                                                                                                                                                                                                                                                                           2 NATFYFKIDNVKKAR 16
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             COUNTRY: USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MOLECULE TYPE:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     RESULT 4
US-08-911-321-3
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Patent No. 6143719
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: The Regents of the University of Michigan
APPLICANT: Schmaler, Alvin H.
APPLICANT: Hasan, Ahmed A.K.
TITLE OF INVENTION: Bradykinin Analogs As Selective Thrombin Inhibitors
FILE REFERENCE: 8820-2 US
CURRENT APPLICATION UNABER: US/08/676,242C
CURRENT FILING DATE: 2000-07-16
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Gaps
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                                                                                                                                                                                                              APPLICANT: Schmaler, Alvin H.
APPLICANT: Jiang, Yongping
TITLE OF INVENTION: Modulation of Blood
TITLE OF INVENTION: Pressure by Altering Bradykinin Levels
CORRESPONDENCE: 2
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 117;
                                                                                                                                                                                                                                                                                                                                                   ADDRESSEE: Temple University - Of the ADDRESSEE: Commonwealth System of Higher Education STREET: 406 University Services STREET: Building
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 100.0%; Score 84; DB 5; 100.0%; Pred. No. 5.3e-07;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ZIP: 19122
COMPUTER READBALE FORM:
MEDIUM TYPE: Diskette, 3.50 inch, 720 Kb
COMPUTER: IBM PS/2
OPERATING SYSTEM: MS-DOS
SOFTWARE: WordPerfect 5.1
CURRENT APPLICATION DATA:
APPLICATION UNDER: PCT/US92/06809
FILING DATE: 19910813
CLASSIFICATION: 514
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PRIOR APPLICATION DATA:
APPLICATION NUMBER: U.S. Application
APPLICATION NUMBER: Serial No. 744,545
FILING DATE: 13 August 1991
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            NAME: MODACO, DANIEL A.
REGISTRATION NUMBER: 30,480
REFERENCE/DOCKET NUMBER: 6056-137
TELECOMMUNICATION INFORMATION:
                                                                                                                                                                       Sequence 1, Application PC/TUS9206809 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TELEPHONE: (215) 568-8383
TELEFAX: (215) 568-5549
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     INFORMATION FOR SEQ ID NO: 1:
SEQUENCE CHARACTERISTICS:
LENGTH: 117 amino acids
TYPE: AMINO ACID
                                     30 NNATFYFKIDNVKKAR 45
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match 100.
Best Local Similarity 100.
Matches 16; Conservative
                1 NNATFYFKIDNVKKAR 16
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                                                                                                                                                                                                                                                                                                                                                                                                                                      CITY: Philadelphia
STATE: Pennsylvania
COUNTRY: U.S.A.
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                                                                                                                             RESULT 2
PCT-US92-06809-1
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US-08-676-242-15
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Patent No. 6303295
GENERAL INFORMATION:
APPLICANT: Taylor, Ethan W.
TITLE OF INVENTION: SELENOPROTEINS, CODING SEQUENCES AND METHODS
FILE REFERENCE: 55-95
CURRENT APPLICATION NUMBER: US/08/679,493A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 872;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CLASSIFICATION: 435
PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA:
APPLICATION DATA: described below:
APPLICATION NUMBER: 60/008,688
FILING DATE: December 15, 1995
ATTORNEY/AGENT INFORMATION:
NAME: Warburg, Richard J.
REGISTRATION NUMBER: 32,327
                                                                                                                                                                                                                                                                                                                        APPLICANT: Mamone, Joseph A.
APPLICANT: Davis, Maria
APPLICANT: Sha, Dan
TITLE OF INVENTION: THERMOSTABLE DNA POLYMERASE
NUMBER OF SEQUENCES: 28
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 38; DB 1; 1
Pred. No. 1.9e+02;
2; Mismatches 5;
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MEDIUM TYPE: 3.5" Diskette, 1.44 Mb
MEDIUM TYPE: 3.5" Medium Type: 1.44 Mb
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MEDIUM TYPE: 3.5" Mb
MEDIUM TYPE: 3.5
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TELECOMONICATION INFORMATION:
TELEPAX: (213) 489-1600
TELEFAX: (213) 955-0440
TELEEX: (7-3510
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADDRESSEE: Lyon & Lyon
STREET: 633 West Fifth Street
STREET: Suite 4700
                                                                                                                                                                                                                                    Sequence 2, Application US/08766014 Patent No. 5744312 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    45.2%;
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Best Local Similarity 50.0
Matches 7; Conservative
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                                 1 NNATFYFKIDNVKK 14
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                                                                                         8 NKLEFFSLIDNIKK 21
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  STREET: Suite 4700 CITY: Los Angeles STATE: California COUNTRY: U.S.A. ZIP: 90071-2066
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TYPE: amino acid
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                                                                                                                                                                             RESULT 6
US-08-766-014-2
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                                                                                                                                                                                                                                                                                                                        Score 40; DB 3; Length 179;
Pred. No. 17;
2; Mismatches 3; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Sequence 3. Application US/08766014
; Patent No. 5744312
; GENERAL INFORMATION:
APPLICANT: Mamone, Joseph A.
APPLICANT: Davis, Maria
APPLICANT: Sha, Dan
TITLE OF INVENTION: THERMOSTABLE DNA POLYMERASE
NUMBER OF SEQUENCES: 28
CORRESPONDENCE ADDRESS:
ADDRESSEE: Lyon & Lyon
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  PRIOR APPLICATION DATA:
PRIOR APPLICATION DATA: including application
PRIOR APPLICATION DATA: described below:
APPLICATION NUMBER: 60/008,688
FILING DATE: December 15, 1995
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                   ; LOCATION:
; IDENTIFICATION METHOD: Deduced sequence
; OTHER INFORMATION: ICP18.5
US-08-911-321-3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               2; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     COUNTRY: U.S.A.
ZIP: 90071-2066
COMPUTER READABLE FORM:
MEDIUM TYPE: 3.5" Diskette, 1.44 Mb
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            COMPUTER: IBM Compatible
OPERATING SYSTEM: IBM P.C. DOS 5.0
SOFTWARE: FastSeq Version 2.0
CURRENT APPLICATION DATA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              APPLICATION NUMBER: US/08/766,014 FILING DATE: Herewith CLASSIFICATION: 435
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     NAME: Warburg, Richard J.
REGISTRATION NUMBER: 32,327
REFERENCE/DOCKET NUMBER: 223/104
TELECOMMUNICATION INFORMATION:
Feline herpesvirus-1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADDRESSEE: Lyon & Lyon
STREET: 633 West Fifth Street
STREET: Suite 4700
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (213) 489-1600
(213) 955-0440
7-3510
                                                       INDIVIDUAL ISOLATE: C-27
CELL TYPE: N/A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            LENGTH: 608 amino acids TYPE: amino acid
                                                                                                                                                                                                                                                                                                        Ouery Match
Best Local Similarity 54.5
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Best Local Similarity 50.0°
Matches 7; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   TELEFAX: (213) 955-044
TELEX: 67-3510
INFORMATION FOR SEQ ID NO:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CITY: Los Angeles
STATE: California
                                                                                                                                                                                                                                                                                                                                                                                                                                                2 NATFYFKIDNV 12
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  linear
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       STRANDEDNESS:
TOPOLOGY: lin
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TELEPHONE:
ORGANISM:
                                                                                                                                                 NAME/KEY:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 5
US-08-766-014-3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       US-08-766-014-3
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Gaps

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US-09-282-305-12

US-09-282-305-12

Sequence 12, Application US/09282305

Patent No. 6287843

SEMERAL INFORMATION:

APPLICANT: Baldwin, Donald A.

APPLICANT: Briggs, Steven P.

APPLICANT: Crane, Virginia C.

TITLE OF INVENTION: Maize Histone Deacetylases And Their Uses

FILE REFERENCE: 5718-44

CURRENT APPLICATION NUMBER: US/09/282,305

CURRENT FILING DATE: 1999-03-31

PRIOR FILING DATE: 1999-04-03

NUMBER OF SEQ ID NOS: 18
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: GERLACH, GERALD F.
APPLICANT: WILLSON, PHILIP J.
APPLICANT: ROSSI-CAMPOS, AMALIA
APPLICANT: POTTER, AMALIA
TITLE OF INVENTION: ACTINOBACILLUS PLEUROPNEUMONIAE OUTER
TITLE OF INVENTION: MEMBRANE LIPOPROTEIN A AND USES THEREOF
CORRESPONDENCE ADDRESS:
ADDRESSEE: GO MUDDLEFIELD ROAD, SUITE 200
CITY: MENLO PARK
                                                     Score 37; DB 4; Length 302;
Pred. No. 93;
2; Mismatches 5; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 37; DB 4; Length 305;
Pred. No. 94;
2; Mismatches 5; Indels
                                                                                                 Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             COMPUTER READABLE FORM:

WEDIUM TYPE: Floppy disk
COMPUTER: IBM PC COMPATIBLE
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.30
CURRENT APPLICATION NDATA:
APPLICATION NUMBER: US/08/515,251A
FILING DATE: 15-AUG-1995
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          JMBER: US 07/971,558
05-NOV-1992
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; Patent No. 5891677
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          44.08;
53.38;
                                                     44.0%;
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                               Query Match
Best Local Similarity 53.37
Best Local 8, Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity 53.3
Matches 8; Conservative
                                                                                                                                   1 NNATFYFKIDNVKKA 15
                                                                                                                                                                         38 DNALMYVKIDDOKLA 52
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APPLICATION NUMBER: U
FILING DATE: 05-NOV-1
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; ORGANISM: Zea mays
US-09-282-305-12
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CLASSIFICATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         COUNTRY: U
            US-09-282-305-14
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SEQ ID NO 12
LENGTH: 305
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Batent No. 6380370
GENERAL INFORMATION:
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: LYNN DOUGETEG-ACID AMINO ACID SEQUENCES RELATING TO STAPHYLOCOCCUS TITLE OF INVENTION:
ELIB REFERENCE: GTC-007
CURRENT APPLICATION NUMBER: US/09/134,001C
CURRENT APPLICATION NUMBER: US 60/064,964
PRIOR PAPLICATION NUMBER: US 60/064,964
PRIOR PELING DATE: 1997-11-08
PRIOR PELING DATE: 1997-11-08
PRIOR PELING DATE: 1997-08-14
NUMBER FO SEQ ID NOS: 5674
SEQ ID NO 3050
LENGTH: 229
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Patent No. 6287843

GENERAL INFORMATION:
APPLICANT: Baldwin, Donald A.
APPLICANT: Brigs, Steven P.
APPLICANT: Crane, Virginia C.
TITLE OF INVENTION: Maize Histone Deacetylases And Their Uses
FILE REPRENCE: 5718-44,
CURRENT APPLICATION NUMBER: US/09/282,305

CURRENT FILING DATE: 1999-03-31

PRIOR APPLICATION NUMBER: 60/080,563

PRIOR FILING DATE: 1999-04-03

NUMBER OF SEQ ID NOS: 18

SEOFTWARE: Patentin Ver. 2.0
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                                                                                                                                                                                                                                                                                                                 1; Mismatches
CURRENT FILING DATE: 1996-07-12
PRIOR APPLICATION NUMBER: 60/001203
PRIOR FILING DATE: 1995-07-14
PRIOR APPLICATION NUMBER: 60/003,112
PRIOR FILING DATE: 1995-09-01
NUMBER OF SEQ ID NOS: 216
SOFTWARE: Patentin Ver. 2.0
LENGTH: 149
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ; TYPE: PRT
; ORGANISM: Staphylococcus epidermidis
US-09-134-001C-3050
                                                                                                                                                                                                                                                                           44.08;
                                                                                                                                                                                                           ; ORGANISM: Escherichia coli
US-08-679-493A-85
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Ouery Match
Best Local Similarity 87.5'
Matches 7; Conservative
                                                                                                                                                                                                                                                                                            Best_Local Similarity 63.6
Matches 7; Conservative
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| 174 FYFKEDNV 181
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ORGANISM: Zea mays
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5 FYFKIDNV 12
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                                                                                                                                                                                          TYPE: PRT
                                                                                                                                                                                                                                                                             Query Match
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Gaps

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NUMBER OF SEQUENCES: 85
CORRESPONDENCE ADDRESS:
ADDRESSEE: No. 56865930 No. 5686593disk of No. 5686593th America, Inc.
                                                                                                 Gaps
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2
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APPLICANT: Woeldike, Helle F.
APPLICANT: Hagen, Frederick
APPLICANT: HJort, Carsten M.
APPLICANT: Sven, Hastrup
TITLE OF INVENTION: An Enzyme Capable of Degrading Cellulose
TITLE OF INVENTION: An Enzyme
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 514;
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COUTRY: United States of America
COUTRY: United States of America
CONTRY: United States of America
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
COMPUTER: 07-UN-1995
FILING DATE: 07-UN-1994
APPLICATION NUMBER: US 07/940,860
FILING DATE: 28-CT-1992
APPLICATION NUMBER: US 07/940,860
FILING DATE: 28-CT-1992
APPLICATION NUMBER: DK 1158/90
FILING DATE: 09-MAY-1991
APPLICATION NUMBER: DK 1158/90
FILING DATE: 08-MAY-1991
APPLICATION NUMBER: DR 1150-1
                                                 DB 1;
                                               Score 36.5; DB 1
Pred. No. 2e+02;
0; Mismatches
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Pred. No. 2e+02
0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              405 Lexington Avenue, 62nd Floor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        NAME: Lambiris, Elias J.
REGISTRATION NUMBER: 33,728
REFERENCE/DOCKET NUMBER: 3435.204-US
                                                                                                                                                                                                                                                                                                      Sequence 21, Application US/08479939 Patent No. 5686593
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TELECOMMUNICATION INFORMATION:
TELEPHONE: 212-667-013
TELEPAX: 212-667-0298
INFORMATION FOR SEQ ID NO: 21:
                                                 43.5%;
                                                                                                                                                                                    287 NKTFYGPGSNFNIDTTKK 304
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                                                                                                                                            2 NATFY----FKIDNVKK 14
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQUENCE CHARACTERISTICS:
LENGTH: 514 amino acids
TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            2 NATFY----FKIDNVKK 14
                                      Query Match
Best Local Similarity 50.0%
Matches 9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ; MOLECULE TYPE: protein US-08-479-939-21
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
Matches 9; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              linear
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    US-08-361-920-21
                                                                                                                                                                                                                                                                                 US-08-479-939-21
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              STREET:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              δ
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CORRESPONDENCE ADDRESS:
ADDRESSEE: No. 54570460 No. 5457046disk of No. 5457046th America, Inc.
STREET: 405 Lexington Avenue, 62nd Floor
                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                                                                 ö
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sequence 11, Application US/08361920
Patent No. 5457046
GENERAL INFORMATION:
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: Hooth, Helle F.
APPLICANT: Hjort, Carsten M.
APPLICANT: Sven, Hastrup M.
APPLICANT: Sven, Hastrup M.
APPLICANT: Sven, Hastrup M.
APPLICANT: OF INVENTION: An Enzyme Capable of Degrading Cellulose
TITLE OF INVENTION: Or Hemicellulose
                                                                                                                                                                                                                                                                                                                                                 Length 365;
                                                                                                                                                                                                                                                                                                                                                                                              3; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ZIP: 10174-6201
COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/08/361,920
                                                                                                                                                                                                                                                                                                                                            Score 37; DB 2; ; Pred. No. 1.1e+02; 0; Mismatches 3:
                                                                    9001-0027.10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   FILING DATE:
CLASSIFICATION: 435
PRIOR APPLICATION 1435
PRICE APPLICATION NUMBER: US 07/940,860
FILING DATE: 28-OCT 1992
APPLICATION NUMBER: DK 1158/90
FILING DATE: 09-MAY-1990
PRIOR APPLICATION NUMBER: PCT/DK91/00124
APPLICATION NUMBER: PCT/DK91/00124
FILING DATE: 08-MAY-1991
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          3435.204-US
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                                 REGISTRATION NUMBER: 33,208
REFERENCE/DOCKET NUMBER: 9001-
TELECOMMUTCATION INFORMATION:
TELEPHONE: (650) 325-7812
TELEPHONE: (650) 325-7823
INFORMATION FOR SEQ ID NO: 2:
SEQUENCE CHARACTERISTICS:
LENGTH: 365 amino acids
TYPE: amino acids
TYPE: amino acids
TYPE: amino acids
TOPOLOGY: linear
MOLECULE TYPE: protein
US-08-515-251a-2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     NAME: Lambiris, Elias J.
REGISTRATION NUMBER: 33,728
REFERENCE/POCKET NUMBER: 3435
TELECOMMUNICATION INFORMATION:
TELEPHONE: 212-867-0123
TELEFAX: 212-867-0298
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               INFORMATION FOR SEQ ID NO: 21: SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                                                                                 44.08;
70.08;
ATTORNEY/AGENT INFORMATION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            : 514 amino acids
amino acid
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Best Local Similarity 70.0°
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TOPOLOGY: linear
MOLECULE TYPE: protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   309 NGTLYSKIDN 318
                                                                                                                                                                                                                                                                                                                                                                                                                                         2 NATFYFKIDN 11
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CITY: New York STATE: New York
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   COUNTRY:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 12
US-08-361-920-21
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                                                                                                                                                                                                                                                       ADDRESSEE No. 57632540 No. 5763254disk of No. 5763254th America, Inc.
STREET: 405 Lexington Avenue, 62nd Floor
CITY: New York
STATE: New York
CONTRY: United States of America
ZIP: 10174-6201
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
COMPUTER: PatentIn Release #1.0, Version #1.25
CURRENT APPLICATION NUMBER: US/08/483,432
FILING DATE: 07-UNI-1995
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          5
                                                                                           APPLICANT: Hagen, Frederick
APPLICANT: Hjort, Carsten M.
APPLICANT: Sven, Hastrup
TITLE OF INVENTION: An Enzyme Capable of Degrading Cellulose
TITLE OF INVENTION: or Hemicellulose
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 36.5; DB 1; Length 514;
Pred. No. 2e+02;
0; Mismatches 4; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                FILING DATE:
APPLICATION NUMBER: US 07/940,860
FILING DATE: 28-0CT-1992
APPLICATION NUMBER: DK 1158/90
FILING DATE: 09-MAY-1990
PRIOR APPLICATION DATE:
APPLICATION NUMBER: PCT/DK91/00124
FILING DATE: 08-MAY-1991
ATTORNEY/AGENT INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  NAME: Lambiris, Elias J.
RECISTRATION NUMBER: 33,728
REFERNCE/DOCKET NUMBER: 3435.204-US
TELECOMMUNICATION INFORMATION:
TELEPHONE: 212-867-0123
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        APPLICANT: Greiner, Steffen
APPLICANT: Universit, t Heidelberg
APPLICANT: Rausch, Thomas
APPLICANT: Krausgrill, Silke
TITLE OF INVENTION: Invertase-Inhibitor
FILE REFERENCE: 25048-13
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       FILING DATE: 07-JUN-1995
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: US/08/361,920
Sequence 21, Application US/08483432 Patent No. 5763254
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Sequence 5, Application US/09230670C Patent No. 6384300 GENERAL INFORMATION:
                                              GENERAL INFORMATION:
APPLICANT: Woeldike, Helle F.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               INFORMATION FOR SEQ ID NO: 21:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity 50.0%;
Matches 9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         287 NKTFYGPGSNFNIDTTKK 304
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 LENGTH: 514 amino acids TYPE: amino acid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2 NATFY-----FKIDNVKK 14
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TELEFAX: 212-867-0298
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ; TOPOLOGY: linear
; MOLECULE TYPE: protein
US-08-483-432-21
                                                                                                                                                                                                               NUMBER OF SEQUENCES: 8
CORRESPONDENCE ADDRESS:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               US-09-230-670C-5
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; CURRENT APPLICATION NUMBER: US/09/230,670C
; CURRENT FILING DATE: 1998-04-05
; PRIOR APPLICATION NUMBER: PG7/EP97/04153
; PRIOR APPLICATION NUMBER: PG7/EP97/04153
; PRIOR FILING DATE: 1998-06-05
; NUMBER OF SEQ ID NOS: 9
; SCOTWARE: Parentin Ver. 2.1
; SEQ ID NO 5
; LENGTH: 159
; TYPE: PRIOR APPLICATION APPLICATION OF TO 70; Pred. NO. ```

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Sequence 6055, Application US/09738626
Publication No. US20020197605A1
GENERAL INFORMATION:
APPLICANT: NAKAGAMA, SATOSHI
APPLICANT: MIXOGUCHI, HIROSHI
APPLICANT: ANDO, SEIKO
APPLICANT: HAYASHI, MIKIRO
  NUMBER OF SEQ ID NOS: 73
SOFTWARE: PatentIn version 3.1
   Query Match 67.9
Best Local Similarity 100.
Matches 11; Conservative
  ) ORGANISM: Homo sapiens
US-09-935-682-3
  YFKIDNVKKAR 16
   1 YFKIDNVKKAR 11
  US-09-738-626-6055
   SEQ ID NO 3
   셤
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  Sequence 3, Appli
Sequence 6055, Ap
Sequence 10580, A
Sequence 10580, A
Sequence 14, Appl
Sequence 12, Appl
Sequence 13653, A
Sequence 13653, Ap
Sequence 126, App
Sequence 129, App
   Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
  Sequence 129, A   March 13, 2003, 19:12:07 ; Search time 14 Seconds
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GenCore version 5.1.4_p5_4578 Copyright (c) 1993 - 2003 Compugen Ltd.
  0.50-09-738-626-6055

0.50-09-815-242-4973

0.50-09-815-242-10580

0.50-09-813-720-14

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US-09-989-736-129
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US-09-990-436-129
US-09-991-181-129
US-09-993-687-129
  Total number of hits satisfying chosen parameters:
  US-09-935-682-3
  199416 seqs, 46092074 residues
   SUMMARIES
  Published_Applications_AA:*
  Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
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Gapop 10.0 , Gapext 0.5
  1 NNATFYFKIDNVKKAR 16
   US-09-461-061A-10
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length: 2000000000
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Match Length
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   Title:
Perfect score:
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Maximum DB s
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   Sequence:
  Searched:
  Database
   Run on:
   Result
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Gaps

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67.9%; Score 57; DB 10; Length 16; 100.0%; Pred. No. 0.0021; ive 0; Mismatches 0; Indels

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Gaps
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   Sequence 40, Application US/10010901
Fatent No. US20020098201A1
GENERAL INFORMATION:
APPLICANT: MCFAdden, Grant
TILLE OF INVENTION: NOVEL MYXOMA GENES FOR IMMUNE MODULATION
FILE REPRENCE: 50082/09002
CURRENT FILING DATE: 2001-12-06
PRIOR PILING DATE: 2001-12-06
PRIOR FILING DATE: 2000-07-12
NUMBER OF SEQ ID NOS: 40
SOFTWARE: FastSEQ for Windows Version 4.0
  Query Match 45.2%; Score 38; DB 10; Length 586; Best Local Similarity 43.8%; Pred. No. 1e+02; Matches 7; Conservative 3; Mismatches 6; Indels
    45.2%; Score 38; DB 10; Length 585; 43.8%; Pred. No. 1e+02;
  APPLICANT: Carr, Grant J.
APPLICANT: Carr, Grant J.
APPLICANT: Carr, Grant J.
APPLICANT: Tamamoto, Robert T.
APPLICANT: Yamamoto, Robert T.
TILLE OF INVENTION: Identification of Essential Genes in TITLE OF INVENTION: Prokaryotes
FILE REFERENCE: ELITRA.011A
CURRENT APPLICATION NUMBER: 00/99/815,242
CURRENT FILING DATE: 2001-03-21
PRIOR FILING DATE: 2000-05-23
PRIOR FILING DATE: 2000-05-23
PRIOR FILING DATE: 2000-05-24
PRIOR FILING DATE: 2000-10-23
PRIOR FILING DATE: 2000-10-23
PRIOR FILING DATE: 2000-11-27
PRIOR FILING DATE: 2000-11-27
PRIOR APPLICATION NUMBER: 60/253,625
PRIOR APPLICATION NUMBER: 60/253,625
PRIOR FILING DATE: 2000-11-27
PRIOR APPLICATION NUMBER: 60/257,931
PRIOR FILING DATE: 2000-11-27
  3; Mismatches
  PRIOR APPLICATION NUMBER: 60/269,308
PRIOR FILING DATE: 2001-02-16
NUMBER OF SEQ ID NOS: 14110
SEQ ID NO 10580
LENGTH: 586
LENGTH: 586
   Sequence 10580, Application US/09815242
Pattent No. USZ0020061569Al
GENERAL INFORMATION:
APPLICANT: Haselbeck, Robert
   ; ORGANISM: Enterococcus faecalis US-09-815-242-10580
  APPLICANT: Ohlsen, Kari L.
APPLICANT: Zyskind, Judith W.
APPLICANT: Wall, Daniel
   : Trawick, John D.
Query Match 45.2
Best Local Similarity 43.8
Matches 7; Conservative
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  1 NNATFYFKIDNVKKAR 16
   1 NNATFYFKIDNVKKAR 16
  US-09-815-242-10580
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US-10-010-901-40
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   Gaps
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0
  APPLICANT: Obligation, Kari L.
APPLICANT: Obligation, Kari L.
APPLICANT: Trani, Jouith W.
APPLICANT: Trani, Jouith W.
APPLICANT: Trani, John D.
APPLICANT: Tranick, John D.
APPLICANT: Tranimoto, Robert T.
APPLICANT: Tranimoto, Robert T.
APPLICANT: Tranimoto, Robert T.
APPLICANT: Xu, H. Howard
TITLE OF INVENTION: Identification of Essential Genes in
TITLE OF INVENTION: 1 ELITRA.0112-21
CURRENT FILING DATE: 2001-03-21
PRIOR APPLICATION NUMBER: 60/296,848
PRIOR APPLICATION NUMBER: 60/206,848
PRIOR PILING DATE: 2000-05-26
PRIOR FILING DATE: 2000-05-26
PRIOR FILING DATE: 2000-05-26
PRIOR FILING DATE: 2000-10-23
PRIOR FILING DATE: 2000-11-27
PRIOR FILING DATE: 2000-11-22
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PRIOR FILING DATE: 2000-11-22
PRIOR FILING DATE: 2000-12-22
PRIOR FILING DATE: 2001-12-22
PRIOR APPLICATION NUMBER: 60/259,308
PRIOR FILING DATE: 2001-12-22
PRIOR FILING DATE: 2001-21-10
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  Score 38; DB 9; Length 282;
Pred. No. 50;
3; Mismatches 0; Indels
   APPLICANT: TATEISHI, NACKO
APPLICANT: SENOH, AKHHIRO
APPLICANT: IKEDA, MASATO
APPLICANT: IKEDA, MASATO
TITLE OF INVENTION: NOVEL POLYNUCLEOTIDES
FILE REFERENCE: 249-125
CURRENT APPLICATION NUMBER: US/09/738,626
CURRENT FILING DATE: 2000-12-18
PRIOR PELING DATE: 1999-12-16
PRIOR FILING DATE: 1999-12-16
PRIOR FILING DATE: 1999-12-16
PRIOR FILING DATE: 2000-04-07
PRIOR FILING DATE: 2000-04-07
PRIOR FILING DATE: 2000-04-07
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SPRIOR FILING DATE: 2000-04-07
SOFTWARE: PARENT PAREN
   Sequence 4973, Application US/09815242
patent No. US20020061569A1
GENERAL INFRMATION:
APPLICANT: Haselbeck, Robert
   ; ORGANISM: Corynebacterium glutamicum US-09-738-626-6055
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Best Local Similarity 66...
  |||:|:|:|
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   RESULT 3
US-09-815-242-4973
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APPLICANT: Summers, Eric
TITLE OF INVENTION: Methods for Improving Secondary Metabolite Production in Fungi
FILE REFERENCE: 109272.147
   ö
  Gaps
   ö
  Score 37; DB 10; Length 359;
Pred. No. 92;
   APPLICANT: Call, Glandouto, Robert T.
APPLICANT: Yanmanto, Robert T.
APPLICANT: Xu, H. Howard
TITLE OF INVENTION: Identification of Essential Genes in
TITLE OF INVENTION: Prokaryotes
FILE REFERENCE: ELITRA.011A
CURRENT APPLICATION NUMBER: US/09/815,242
CURRENT FILING DATE: 2000-03-21
PRIOR APPLICATION NUMBER: 60/191,078
PRIOR FILING DATE: 2000-05-23
PRIOR FILING DATE: 2000-05-26
PRIOR FILING DATE: 2000-10-23
PRIOR FILING DATE: 2000-11-27
PRIOR FILING DATE: 2001-10-216
NUMBER OF SEQ ID NOS: 14110
SEQ ID NO 13653
FROM FILING DATE: 2001-02-16
NUMBER OF SEQ ID NOS: 14110
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  Maxon, Mary
Milne, Todd
No. US20020128250Alman, Thea
   Sequence 13653, Application US/09815242 Patent No. US20020061569A1
   Sequence 376, Application US/09801368 Patent No. US20020128250A1 GENERAL INFORMATION:
   ; ORGANISM: Streptococcus pneumoniae US-09-815-242-13653
   APPLICANT: Haselbeck, Robert
APPLICANT: Ohlsen, Kari L.
APPLICANT: 2yskind, Judith W.
APPLICANT: Wall, Daniel
  44.0%;
ilarity 50.0%;
Conservative
   APPLICANT: Trawick, John D. APPLICANT: Carr, Grant J.
  APPLICANT: Busby, Robert
APPLICANT: Call, Brian
APPLICANT: Hecht, Peter
APPLICANT: Holtzman, Doug
APPLICANT: Madden, Kevin
  Salama, Sofie
            38 DNALMYVKIDDOKLA 52
  Sherman, Amir
   324 THYFKLSEIEKA 335
  Royer, John
  4 TFYFKIDNVKKA 15
   Silva, Jefi
   Query Match
Best Local Similarity
Matches 6; Conserv
   US-09-801-368-376
  APPLICANT:
APPLICANT:
APPLICANT:
   APPLICANT:
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   Gaps
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  Gaps
  uS-09-883-720-12

iSequence 12, Application US/09883720

iSequence 12, Application US/09883720

iSequence 12, Application US/09883720

iSequence 12, Application US/09883720

iSeneral INFORMATION:

APPLICANT: Briggs, Steven P.

iAPPLICANT: Crane, Virginia C.

iTILE REFERENCE: 5718-44,

iCURRENT FILING NUMBER: 09/282, 305

PRIOR APPLICATION NUMBER: 09/282, 305

PRIOR PELING DATE: 1999-03-31

iNUMBER OF EQ ID NOS: 18

iSEQ ID NO 12

iENGTH: 305

municipal and partical and
   Sequence 14, Application US/09883720

Patent No. US200202256a1

GENERAL INFORMATION:

APPLICANT: Baldwin, Donald A.

APPLICANT: Bridgs, Steven P.

APPLICANT: Crane, Virginia C.

TITLE OF INVENTION: Malze Histone Deacetylases And Their Uses

FILE REFERENCE: 5718 44,

CURRENT APPLICATION NUMBER: US/09/883,720

CURRENT APPLICATION NUMBER: 09/282,305

PRIOR FILING DATE: 1999-03-31

NUMBER OF SEQ ID NOS: 18

SOFTWARE: Patentin Ver. 2.0
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  ö
   DB 10; Length 302; 78;
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44.0%; Score 37; DB 10; Length 305;
Best Local Similarity 53.3%; Pred. No. 78;
Matches 8; Conservative 2; Mismatches 5; Indels
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57.1%;
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  1 NNATFYFKIDNVKKA 15
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US-10-010-901-40
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; ORGANISM: Zea mays
US-09-883-720-14
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; ORGANISM: Zea mays
US-09-883-720-12
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US-09-883-720-14
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LENGTH: 302
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TITLE OF INVENTION: Secreted and Transmembrane Polypeptides and Nucleic TITLE OF INVENTION: Acids Encoding the Same FILE REFERENCE: P2730PIC20 CURRENT APPLICATION NUMBER: US/09/992,598 CURRENT FILING DATE: 2001-11-14 PRIOR APPLICATION NUMBER: 60/049787
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                      PRIOR APPLICATION NUMBER: 60/242,578
PRIOR APPLICATION NUMBER: 60/242,578
PRIOR FILING DATE: 2000-10-23
PRIOR FILING DATE: 2000-11-27
PRIOR FILING DATE: 2000-11-27
PRIOR FILING DATE: 2000-12-22
PRIOR FILING DATE: 2000-12-22
PRIOR FILING DATE: 2001-02-16
NUMBER OF SEQ ID NOS: 14110
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SOFTWARRE: FASTESEQ for Windows Version 4.0
LENGTH: 400
  ; LOCATION: (1)...(400)
; OTHER INFORMATION: Xaa - Any Amino Acid
US-09-815-242-5324
   Sequence 129, Application US/09992598 Patent No. US20020160384A1 GENERAL INFORMATION:
  PRIOR FILING DATE: 1997-10-17
PRIOR APPLICATION NUMBER: 60/065186
PRIOR FILING DATE: 1997-11-12
PRIOR APPLICATION NUMBER: 60/065311
PRIOR FILING DATE: 1997-11-13
PRIOR APPLICATION NUMBER: 60/066770
  PRIOR FILING DATE: 1997-06-16
PRIOR APPLICATION NUMBER: 60/062250
  Godowski, Paul J.
Grimaldi, J.Christopher
   TYPE: PRT ORGANISM: Staphylococcus aureus
   Watanabe, Colin K.
Williams, P. Mickey
Wood, William I.
   Stewart, Timothy A
           PRIOR FILING DATE: 2000-05-26
  Ferrara, Napoleone
  Gerritsen, Mary E.
Goddard, Audrey
   Pan, James
Paoni, Nicholas F.
   Roy, Margaret Ann
  Gerber, Hanspeter
   Gurney, Austin L.
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   Kljavin, Ivar J.
  APPLICANT: Ashkenazi, Avi J.
  Botstein, David
   Napier, Mary A.
  Desnoyers, Luc
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   Tumas, Daniel
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  Eaton, Dan L.
  Zhang, Zemin
  Query Match
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  NAME/KEY: VARIANT
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  GENERAL INFORMATION:
APPLICANT: Rosen et al.
TITLE OF INVENTION: Nucleic Acids, Proteins, and Antibodies
FILE REPERENCE: PCOOS
CURRENT APPLICATION NUMBER: US/09/764,877
CURRENT FILING DATE: 2001-01-17
Prior application data removed - refer to PALM or file wrapper NUMBER OF SEQ ID NOS: 4031
SOFTWARE: PatentIn Ver. 2.0
   Score 37; DB 10; Length 717;
Pred. No. 1.9e+02;
  42.9%; Score 36; DB 10; Length 133; 42.1%; Pred. No. 50;
  APPLICANT: Object.
APPLICANT: Object.
APPLICANT: Syskind, Judith W.
APPLICANT: Zyskind, Judith W.
APPLICANT: Wall, Daniel
APPLICANT: Tradick, John D.
APPLICANT: Tradick, John D.
APPLICANT: Tamamoto, Robert T.
APPLICANT: Xu, H. Howard
TITLE OF INVENTION: Dentification of Essential Genes in
TITLE OF INVENTION: Prokaryotes
TITLE OF INVENTION: 1040-11A
CURRENT APPLICATION NUMBER: US/09/815,242
CURRENT APPLICATION NUMBER: 60/191,078
PRIOR FILING DATE: 2000-03-21
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CURRENT FILING DATE: 2001-03-07
PRIOR APPLICATION NUMBER: US 09/487,558
PRIOR FILING DATE: 2000-01-19
PRIOR FILING DATE: 1999-10-20
NUMBER OF SEQ ID NOS: 440
SOFTWARE: Patentin version 3.0
LENGTH: 717
  Sequence 1463, Application US/09764877 Patent No. US20020147140A1
  Sequence 5324, Application US/09815242
Patent No. US20020061569A1
GENERAL INFORMATION:
APPLICANT: Haselbeck, Robert
   TYPE: PRT CRGANISM: Saccharomyces cerevisiae US-09-801-368-376
   44.0%;
  2 NATFYFKID----NVKKAR 16
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| 997-11-24 MBER: 60/075945 998-02-25 998-03-20 998-03-20 MBER: 60/08322 98-04-28 MBER: 60/084600 998-05-07 998-05-07 MBER: 60/087106 998-05-28 MBER: 60/087106 998-05-07 MBER: 60/087106 998-05-07 MBER: 60/087607               |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | 8 - 0 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1                                                                                                                               |
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PRIOR APPLICATION NUMBER: 60,08538

PRIOR APPLICATION NUMBER: 60,08559

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PRIOR PRILIKG DATE: 1998-06-17

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PRIOR PLILIKG DATE: 1998-06-18

PRIOR PAPLICATION NUMBER: 60,08901

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FILING DATE: 1998-05-28
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  FILING DATE: 1998-06-02
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APPLICATION NUMBER: 60/087827
   APPLICATION NUMBER: 60/088021
FILING DATE: 1998-06-04
   APPLICATION NUMBER: 60/088025
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  APPLICATION NUMBER: 60/088026
   FILING DATE: 1998-06-04
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   APPLICATION NUMBER: 60/089600
  FILING DATE: 1998-06-03
   FILING DATE: 1998-06-04
  1998-05-0
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   APPLICANT: Wood, William I.
APPLICANT: Thang, Zemin
TITLE OF INVENTION: Secreted and Transmembrane Polypeptides and Nucleic
TITLE OF INVENTION: Acids Encoding the Same
FILE REFERENCE: P2730PIC66
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Pred. No. 1.6e+0
  CURRENT APPLICATION NUMBER: US/09/989,293A
CURRENT FILING DATE: 2001-11-20
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PRIOR FILING DATE: 1998-07-02
PRIOR FILING DATE: 1998-07-07
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PRIOR FILING DATE: 1998-07-07
  Sequence 129, Application US/09989293A Patent No. US20020177164A1 GENERAL INFORMATION: APPLICANT: Ashkenazi, Avi J.
  Grimaldi, J. Christopher
Gurney, Austin L.
  Watanabe, Colin K.
Williams, P. Mickey
Wood, William I.
   Paoni, Nicholas F.
Roy, Margaret Ann
Stewart, Timothy A.
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  Gerber, Hanspeter
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  FILING DATE: 1998-04-28
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APPLICANT: Walliams, P. Mickey
APPLICANT: Walliams, P. Mickey
APPLICANT: Walliams, P. Mickey
APPLICANT: Wood, William I.
APPLICANT: Diang, Zemin
TITLE OF INVEWION: Secreted and Transmembrane Polypeptides and Nucleic
TITLE OF INVEWION: Secreted and Transmembrane Polypeptides and Nucleic
TITLE OF INVEWION: Acids Encoding the Same
FILE REFERENCE: P2730PLCG1
CURRENT APPLICATION NUMBER: US/09/989,735
CURRENT FILING DATE: 2001-11-19
PRIOR APPLICATION NUMBER: 60/06250
PRIOR FILING DATE: 1997-11-12
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Publication No. US20020193299A1
GENERAL INFORMATION:
PRIOR APPLICATION NUMBER: 60/091978
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PRIOR APPLICATION NUMBER: 60/092182
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  PRIOR APPLICATION NUMBER: 60/087607 PRIOR FILING DATE: 1998-06-02 PRIOR APPLICATION NUMBER: 60/087609
  Grimaldi, J.Christopher
  Roy, Margaret Ann
Stewart, Timothy A.
Tumas, Daniel
Watanabe, Colin K.
Williams, P. Mickey
Wood, William I.
  Ferrara, Napoleone
  Gerritsen, Mary E.
  Pan, James
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   Gurney, Austin L.
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   239 NMASLYGQLDTTKKLR 254
  APPLICANT: Ashkenazi, Avi J.
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| FILING DATE APPLICATION TITING DATE APPLICATION FILING DATE APPLICATION | PILLING FILLING FILLIN                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            | TON                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| PRIOR                                                                                                                                                                               | PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR<br>PRIOR | PRIOR |

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   APPLICATION NUMBER: 60/089952
   APPLICANT: KOY, MALUGHA AND APPLICANT: Stewart, Timochy A. APPLICANT: Stewart, Timochy A. APPLICANT: Tumas, Daniel APPLICANT: Tumas, Daniel APPLICANT: Tumas, Daniel Watanabe, Collin K. APPLICANT: Watanabe, Collin K. APPLICANT: Walliams, P. Mickey APPLICANT: Williams, P. Mickey APPLICANT: Walliam I. APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANT: APPLICANTON: Sceneted and Transmembrane Polypeptides and Nucleic TITLE OF INVENTION: Acids Encoding the Same FILE OF INVENTION: Acids Encoding the Same FILE OF INVENTION NUMBER: US/09/990,444

PRIOR PILING DATE: 1997-11-13

PRIOR PLILNG DATE: 1997-11-13

PRIOR PPLICATION NUMBER: 60/065186

PRIOR FILING DATE: 1997-11-13

PRIOR PPLICATION NUMBER: 60/065311

PRIOR PLILNG DATE: 1998-11-13

PRIOR PLILNG DATE: 1998-02-25

PRIOR FILING DATE: 1998-02-26

PRIOR FILING DATE: 1998-03-20

PRIOR PLILNG DATE: 1998-04-28

PRIOR PLILNG DATE: 1998-04-28

PRIOR PLILNG DATE: 1998-04-28
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3; Mismatches 6;
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R FILING DATE: 1998-05-07
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FILING DATE: 1998-06-04
  Grimaldi, J. Christopher
Gurney, Austin L.
  Roy, Margaret Ann
Stewart, Timothy A.
Tumas, Daniel
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Conservative
   Fong, Sherman
Gerber, Hanspeter
Gerritsen, Mary E.
   Ferrara, Napoleone
   Pan, James
Paoni, Nicholas F.
  Godowski, Paul J.
  APPLICANT: Ashkenazi, Avi J. APPLICANT: Baker, Kevin P. APPLICANT: Botstein, David
   Kljavin, Ivar J.
  1 NNATEYFKIDNVKKAR 16
   Goddard, Audrey
   Napier, Mary A.
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Best Local Similarity
Matches 7; Conserv
   APPLICANT:
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Search completed: March 13, 2003, 19:15:02 Job time : 16 secs

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R APPLICATION NUMBER: 60/090355
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Matches 7; Conservative

ö

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239 NMASLYGQLDTTKKLR 254